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(54) Title: HUMAN GENOME-DERIVED SINGLE EXON NUCLEIC ACID PROBES USEFUL FOR ANALYSIS OF GENE EXPRESSION IN HUMAN HEAK'I

(57) Abstract: A single exon nucleic acid microarray comprising a plurality of single exon nucleic acid probes for measuring gene expression in a sample derived from human heart is described. Also described are single exon nucleic acid probes expressed in the heart and their use in methods for detecting gene expression.

HUMAN GENOME-DERIVED SINGLE EXON NUCLEIC ACID PROBES USEFUL FOR ANALYSIS OF GENE EXPRESSION IN HUMAN HEART

CROSS REFERENCE TO RELATED APPLICATIONS

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The present application is a continuation-in-part of U.S. patent application serial nos. 09/632,366, filed August 3, 2000 and 09/608,408, filed June 30, 2000; claims the benefit under 35 U.S.C. s 119(e) of U.S.provisional patent application serial nos. 60/236,359, filed September 27, 2000, 60/234,687, filed September 21, 2000, 60/207,456, filed May 26, 2000, and 60/180,312, filed February 4, 2000; and further claims the benefit under 35 U.S.C. s 119(a) of UK patent application no. 0024263.6, filed October 4, 2000, the disclosures of which are incorporated herein by reference in their entireties.

REFERENCE TO SEQUENCE LISTING AND INCORPORATION BY REFERENCE THEREOF

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The present application includes a Sequence Listing in electronic format, filed pursuant to PCT Administrative Instructions 801 - 806 on a single CD-R disc, in triplicate, containing a file named pto_HEART.txt, created 24 January 2001, having 20,186,946 bytes. The Sequence Listing contained in said file on said disc is incorporated herein by reference in its entirety.

Field of the Invention

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The present invention relates to genome-derived single exon microarrays useful for verifying the expression of regions of genomic DNA predicted to encode protein. In particular, the present invention relates to unique genomederived single exon nucleic acid probes expressed in human

heart and single exon nucleic acid microarrays that include such probes.

Background of the Invention

For almost two decades following the invention of general techniques for nucleic acid sequencing, Sanger et al., Proc. Natl. Acad. Sci. USA 70(4):1209-13 (1973); Gilbert et al., Proc. Natl. Acad. Sci. USA 70(12):3581-4 (1973), these techniques were used principally as tools to further the understanding of proteins — known or suspected — about which a basic foundation of biological knowledge had already been built. In many cases, the cloning effort that preceded sequence identification had been both informed and directed by that antecedent biological understanding.

For example, the cloning of the T cell receptor for antigen was predicated upon its known or suspected cell type-specific expression, by its suspected membrane association, and by the predicted assembly of its gene via T cell-specific somatic recombination. Subsequent sequencing efforts at once confirmed and extended understanding of this family of proteins. Hedrick et al., Nature 308(5955):153-8 (1984).

More recently, however, the development of high
throughput sequencing methods and devices, in concert with
large public and private undertakings to sequence the human
and other genomes, has altered this investigational
paradigm: today, sequence information often precedes
understanding of the basic biology of the encoded protein
product.

One of the approaches to large-scale sequencing is predicated upon the proposition that expressed sequences — that is, those accessible through isolation of mRNA — are of greatest initial interest. This "expressed sequence tag" ("EST") approach has already yielded vast

amounts of sequence data (see for example Adams et al.,

Science 252:1651 (1991); Williamson, Drug Discov. Today

4:115 (1999)). For nucleic acids sequenced by this

approach, often the only biological information that is

known a priori with any certainty is the likelihood of

biologic expression itself. By virtue of the species and

tissue from which the mRNA had originally been obtained,

most such sequences are also annotated with the identity of

the species and at least one tissue in which expression

appears likely.

More recently, the pace of genomic sequencing has accelerated dramatically. When genomic DNA serves as the initial substrate for sequencing efforts, expression cannot be presumed; often the only a priori biological information about the sequence includes the species and chromosome (and perhaps chromosomal map location) of origin.

With the ever-accelerating pace of sequence accumulation by directed, EST, and genomic sequencing approaches — and in particular, with the accumulation of sequence information from multiple genera, from multiple species within genera, and from multiple individuals within a species — there is an increasing need for methods that rapidly and effectively permit the functions of nucleic sequences to be elucidated. And as such functional information accumulates, there is a further need for methods of storing such functional information in meaningful and useful relationship to the sequence itself; that is, there is an increasing need for means and apparatus for annotating raw sequence data with known or predicted functional information.

Although the increase in the pace of genomic sequencing is due in large part to technological changes in sequencing strategies and instrumentation, Service, Science 280:995 (1998); Pennisi, Science 283: 1822-1823 (1999), there is an important functional motivation as well.

While it was understood that the EST approach would rarely be able to yield sequence information about the noncoding portions of the genome, it now also appears the EST approach is capable of capturing only a fraction of a genome's actual expression complexity.

For example, when the C. elegans genome was fully sequenced, gene prediction algorithms identified over 19,000 potential genes, of which only 7,000 had been found by EST sequencing. C. elegans Sequencing Consortium, 10 Science 282:2012 (1998). Analogously, the recently completed sequence of chromosome 2 of Arabidopsis predicts over 4000 genes, Lin et al., Nature, 402:761 (1999), of which only about 6% had previously been identified via EST sequencing efforts. Although the human genome has the 15 greatest depth of EST coverage, it is still woefully short of surrendering all of its genes. One recent estimate suggests that the human genome contains more than 146,000 genes, which would at this point leave greater than half of the genes undiscovered. It is now predicted that many 20 genes, perhaps 20 to 50%, will only be found by genomic sequencing.

There is, therefore, a need for methods that permit the functional regions of genomic sequence — and most importantly, but not exclusively, regions that

25 function to encode genes — to be identified.

Much of the coding sequence of the human genome is not homologous to known genes, making detection of open reading frames ("ORFs") and predictions of gene function difficult. Computational methods exist for predicting coding regions in eukaryotic genomes. Gene prediction programs such as GRAIL and GRAIL II, Uberbacher et al., Proc. Natl. Acad. Sci. USA 88(24):11261-5 (1991); Xu et al., Genet. Eng. 16:241-53 (1994); Uberbacher et al., Methods Enzymol. 266:259-81 (1996); GENEFINDER, Solovyev et al., Nucl. Acids. Res. 22:5156-63 (1994); Solovyev et al.,

Ismb 5:294-302 (1997); and GENESCAN, Burge et al., J. Mol.
Biol. 268:78-94 (1997), predict many putative genes without
known homology or function. Such programs are known,
however, to give high false positive rates. Burset et al.,
Genomics 34:353-367 (1996). Using a consensus obtained by
a plurality of such programs is known to increase the
reliability of calling exons from genomic sequence.
Ansari-Lari et al., Genome Res. 8(1):29-40 (1998)

Identification of functional genes from genomic
data remains, however, an imperfect art. For example, in
reporting the full sequence of human chromosome 21, the
Chromosome 21 Mapping and Sequencing Consortium reports
that prior bioinformatic estimates of human gene number may
need to be revised substantially downwards. Nature
405:311-199 (2000); Reeves, Nature 405:283-284 (2000).

Thus, there is a need for methods and apparatus that permit the functions of the regions identified bioinformatically — and specifically, that permit the expression of regions predicted to encode protein — readily to be confirmed experimentally.

Recently, the development of nucleic acid microarrays has made possible the automated and highly parallel measurement of gene expression. Reviewed in Schena (ed.), DNA Microarrays: A Practical Approach

25 (Practical Approach Series), Oxford University Press (1999) (ISBN: 0199637768); Nature Genet. 21(1)(suppl):1 - 60 (1999); Schena (ed.), Microarray Biochip: Tools and Technology, Eaton Publishing Company/BioTechniques Books Division (2000) (ISBN: 1881299376).

It is common for microarrays to be derived from cDNA/EST libraries, either from those previously described in the literature, such as those from the I.M.A.G.E. consortium, Lennon et al., Genomics 33(1):151-2 (1996), or from the construction of "problem specific" libraries targeted at a particular biological question, R.S. Thomas

et al., Cancer Res. (in press). Such microarrays by definition can measure expression only of those genes found in EST libraries, and thus have not been useful as probes for genes discovered solely by genomic sequencing.

The utility of using whole genome nucleic acid microarrays to answer certain biological questions has been demonstrated for the yeast Saccharomyces cerevisiae. De Risi et al., Science 278:680 (1997). The vast majority of yeast nuclear genes, approximately 95% however, are single exon genes, i.e., lack introns, Lopez et al., RNA 5:1135-1137 (1999); Goffeau et al., Science 274:563-67 (1996), permitting coding regions more readily to be identified. Whole genome nucleic acid microarrays have not generally been used to probe gene expression from more complex eukaryotic genomes, and in particular from those averaging more than one intron per gene.

Diseases of the heart and vascular system are a significant cause of human morbidity and mortality. Increasingly, genetic factors are being found that contribute to predisposition, onset, and/or aggressiveness of most, if not all, of these diseases. Although mutations in single genes have on occasion been identified as causative, these disorders are for the most part believed to have polygenic etiologies. There is a need for methods and apparatus that permit prediction, diagnosis and prognosis of diseases of the human heart, particularly those diseases with polygenic etiology.

Summary of the Invention

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The present invention solves these and other problems in the art by providing methods and apparatus for predicting, confirming, and displaying functional information derived from genomic sequence. The present invention also provides apparatus for verifying the

expression of putative genes identified within genomic sequence.

In particular, the invention provides novel genome-derived single exon nucleic acid microarrays useful for verifying the expression of putative genes identified within genomic sequence.

The present invention also provides compositions and kits for the ready production of nucleic acids identical in sequence to, or substantially identical in sequence to, probes on the genome-derived single exon microarrays of the present invention.

Accordingly, in a first aspect of the invention, there is provided a spatially-addressable set of single exon nucleic acid probes for measuring gene expression in a sample derived from human heart, comprising a plurality of single exon nucleic acid probes according to any one of the nucleotide sequences set out in SEQ ID NOs: 1 - 9,980 or a complementary sequence, or a portion of such a sequence.

By plurality is meant at least two, suitably at least 20, most suitably at least 100, preferably at least 1000 and, most preferably, upto 5000.

In one embodiment of the first aspect, each of said plurality of probes is separately and addressably amplifiable.

In an alternative embodiment, each of said plurality of probes is separately and addressably isolatable from said plurality.

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In a preferred embodiment, each of said plurality of probes is amplifiable using at least one common primer.

30 Preferably, each of said plurality of probes is amplifiable using a first and a second common primer.

In yet another embodiment, said set of single exon nucleic acid probes comprises between 50 - 20,000 probes, for example, 50 - 5000.

35 Suitably, said set of single exon nucleic acid

probes comprises at least 50 - 1000 discrete single exon nucleic acid probes having a sequence as set out in any of SEQ ID NOS.: 1 - 19,771 or a complimentary sequence, or a portion of such a sequence.

Preferably, the average length of the single exon 5 nucleic acid probes is between 200 and 500 bp. It is preferred that the average length should be at least 200bp, suitably at least 250bp, most suitably at least 300bp, preferably at least 400bp and, most preferably, 500 bp.

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In another embodiment, the single exon nucleic acid probes lack prokaryotic and bacteriophage vector sequence. It is preferred that at least 50%, suitably at least 60%, most suitably at least 70%, preferably at least 75%, more preferably at least 80, 85, 90, 95 or 99% of said 15 single exon nucleic acid probes lack prokaryotic and bacteriophage vector sequence.

In another preferred embodiment, said single exon nucleic acid lack homopolymeric stretches of A or T. It is preferred that at least 50%, suitably at least 60%, most 20 suitably at least 70%, preferably at least 75%, more preferably at least 80, 85, 90, 95 or 99% of said single exon nucleic acid probes lack homopolymeric stretches of A or T.

Preferably, a spatially-addressable set of single exon nucleic acid probes in accordance with the first aspect of the invention is is addressably disposed upon a substrate.

Suitable substrates include a filter membrane which may, preferably, be nitrocellulose or nylon. The 30 nylon may preferably, be positively-charged. Other suitable substrates include glass, amorphous silicon, crystalline silicon, and plastic. Further suitable materials include polymethylacrylic, polyethylene, polypropylene, polyacrylate, polymethylmethacrylate, polyvinylchloride, 35 polytetrafluoroethylene, polystyrene, polycarbonate,

polyacetal, polysulfone, celluloseacetate, cellulosenitrate, nitrocellulose, and mixtures thereof.

In a second aspect of the invention, there is provided a microarray comprising a spatially addressable set of single exon nucleic acid probes in accordance with the first aspect of the invention.

In one embodiment, a genome-derived single-exon microarray is packaged together with such an ordered set of amplifiable probes corresponding to the probes, or one or more subsets of probes, thereon. In alternative embodiments, the ordered set of amplifiable probes is packaged separately from the genome-derived single exon microarray.

In another aspect, the invention provides genomederived single exon nucleic acid probes useful for gene
expression analysis, and particularly for gene expression
analysis by microarray. In particular embodiments of this
aspect, the present invention provides human single-exon
probes that include specifically-hybridizable fragments of
SEQ ID Nos. 9,981 - 19,771, wherein the fragment hybridizes
at high stringency to an expressed human gene. In
particular embodiments, the invention provides single exon
probes comprising SEQ ID Nos. 1 - 9,980.

Accordingly, in a third aspect of the invention,

25 there is provided a single exon nucleic acid probe for
measuring human gene expression in a sample derived from
human heart which is a nucleic acid molecule comprising a
nucleotide sequence as set out in any of SEQ ID NOs.: 1 9,980 or a complementary sequence or a fragment thereof

30 wherein said probe hybridizes at high stringency to a
nucleic acid expressed in the human heart.

In one embodiment, a single exon nucleic acid probe in accordance with the third aspect comprises a nucleotide sequence as set out in any of SEQ ID NOs.: 9,981

35 - 19,771 or a complementary sequence or a fragment thereof.

In a fourth aspect of the invention, there is provided a single exon nucleic acid probe for measuring human gene expression in a sample derived from human heart which is a nucleic acid molecule having a sequence encoding a peptide comprising a peptide sequence as set out in any of SEQ ID NOs.: 19,772 - 29,119 or a complementary sequence or a fragment thereof wherein said probe hybridizes at high stringency to a nucleic acid expressed in the human heart.

Preferably, a single exon nucleic acid probe in accordance with the third or fourth aspects of the invention comprises between at least 15 and 50 contiguous nucleotides of said SEQ ID NO:. It is preferred that the single exon nucleic acid probe comprises at least 15, suitably at least 20, more suitably at least 25 or preferably at least 50 contiguous nucleotides of said SEQ ID NO:.

In another preferred embodiment, a single exon nucleic acid probe in accordance with the third or fourth aspects of the invention is between 3kb and 25kb in length.

20 It is preferred that said probe is no more than 3kb, suitably no more than 5kb, more suitably no more than 10kb, preferably 15kb, more preferably 20kb or, most preferably, no more than 20kb in length.

Preferably, a single exon nucleic acid probe in accordance with either the fifth or sixth aspect of the invention is DNA, preferably single-stranded DNA, RNA or PNA.

In another embodiment of either the third or fourth aspect of the invention, a single exon nucleic acid probe is detectably labeled. Suitable detectable labels include a radionuclide, a fluorescent label or a first member of a specific binding pair. Suitable fluorescent labels include dyes such as cyanine dyes, preferably Cy3 and Cy5 although other suitable dyes will be known to those skilled in the art.

In a particularly preferred embodiment, a single exon nucleic acid probe in accordance with either the third or fourth aspect of the invention lacks prokaryotic and bacteriophage vector sequence. In yet another embodiment, a single exon nucleic acid probe in accordance with either the third or fourth aspect of the invention lacks homopolymeric stretches of A or T.

In a fifth aspect of the invention, there is provided an amplifiable nucleic acid composition, comprising:

the single exon nucleic acid probe in accordance with either of the third or fourth aspects of the invention; and at least one nucleic acid primer;

wherein said at least one primer is sufficient to prime enzymatic amplification of said probe.

In an sixth aspect of the invention, there is provided a method of measuring gene expression in a sample derived from human heart, comprising:

contacting the single exon microarray in

20 accordance with the second aspect of the invention, with a
first collection of detectably labeled nucleic acids, said
first collection of nucleic acids derived from mRNA of
human heart; and then

measuring the label detectably bound to each probe of said microarray.

In a seventh aspect of the invention, there is provided a method of identifying exons in a eukaryotic genome, comprising:

algorithmically predicting at least one exon from 30 genomic sequence of said eukaryote; and then

detecting specific hybridization of detectably labeled nucleic acids to a single exon probe,

wherein said detectably labeled nucleic acids are derived from mRNA from the heart of said eukaryote, said probe is a single exon probe having a fragment identical in

sequence to, or complementary in sequence to, said predicted exon, said probe is included within a single exon microarray in accordance with the first aspect of the invention, and said fragment is selectively hybridizable at 5 high stringency.

In a eighth aspect of the invention, there is provided a method of assigning exons to a single gene, comprising:

identifying a plurality of exons from genomic

10 sequence in accordance with the seventh aspect of the invention; and then

measuring the expression of each of said exons in a plurality of tissues and/or cell types using hybridization to single exon microarrays having a probe

15 with said exon,

wherein a common pattern of expression of said exons in said plurality of tissues and/or cell types indicates that the exons should be assigned to a single gene.

In an ninth aspect of the invention, there is provided a nucleic acid sequence as set out in any of SEQ ID NOs: 1 - 19,771 wherein said sequence encodes a peptide.

In a tenth aspect of the invention, there is provided a peptide encoded by a sequence comprising a

25 sequence as set out in any of SEQ ID NOs: 9,981 - 19,771, or a complementary sequence or coding portion thereof.

In a preferred embodiment, a peptide may be encoded by a sequence comprising a sequence set out in any of SEQ ID NOS.: 1 - 9,980.

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In a further aspect, the invention provides peptides comprising an amino acid sequence translated from the DNA fragments, said amino acid sequences comprising SEQ ID NOS.: 9,981 - 19,771.

Accordingly in a eleventh aspect of the invention there is provided a peptide comprising a sequence as set

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out in any of SEQ ID NOs: 19,772 - 29,119, or fragment thereof.

In another aspect, the invention provides means for displaying annotated sequence, and in particular, for 5 displaying sequence annotated according to the methods and apparatus of the present invention. Further, such display can be used as a preferred graphical user interface for electronic search, query, and analysis of such annotated sequence.

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Detailed Description of the Invention

Definitions

As used herein, the term "microarray" and phrase "nucleic acid microarray" refer to a substrate-bound collection of plural nucleic acids, hybridization to each of the plurality of bound nucleic acids being separately detectable. The substrate can be solid or porous, planar 20 or non-planar, unitary or distributed.

As so defined, the term "microarray" and phrase "nucleic acid microarray" include all the devices so called in Schena (ed.), DNA Microarrays: A Practical Approach (Practical Approach Series), Oxford University Press (1999) 25 (ISBN: 0199637768); Nature Genet. 21(1)(suppl):1 - 60 (1999); and Schena (ed.), Microarray Biochip: Tools and Technology, Eaton Publishing Company/BioTechniques Books Division (2000) (ISBN: 1881299376). As so defined, the term "microarray" and phrase "nucleic acid microarray" 30 further include substrate-bound collections of plural nucleic acids in which the nucleic acids are distributably disposed on a plurality of beads, rather than on a unitary planar substrate, as is described, inter alia, in Brenner et al., Proc. Natl. Acad. Sci. USA 97(4):166501670 (2000); 35 in such case, the term "microarray" and phrase "nucleic

acid microarray" refer to the plurality of beads in aggregate.

As used herein with respect to a nucleic acid microarray, the term "probe" refers to the nucleic acid that is, or is intended to be, bound to the substrate; in such context, the term "target" thus refers to nucleic acid intended to be bound thereto by Watson-Crick complementarity. As used herein with respect to solution phase hybridization, the term "probe" refers to the nucleic acid of known sequence that is detectably labeled.

As used herein, the expression "probe comprising SEQ ID NO.", and variants thereof, intends a nucleic acid probe, at least a portion of which probe has either (i) the sequence directly as given in the referenced SEQ ID NO., or (ii) a sequence complementary to the sequence as given in the referenced SEQ ID NO., the choice as between sequence directly as given and complement thereof dictated by the requirement that the probe hybridize to mRNA.

As used herein, the term "open reading frame" and
the equivalent acronym "ORF" refer to that portion of an
exon that can be translated in its entirety into a sequence
of contiguous amino acids i.e. a nucleic acid sequence
that, in at least one reading frame, does not possess stop
codons; the term does not require that the ORF encode the
entirety of a natural protein.

As used herein, the term "amplicon" refers to a PCR product amplified from human genomic DNA, containing the predicted exon.

As used herein the term "exon" refers to the consensus prediction of the various exon and gene predicting algorithms i.e. a nucleic acid sequence bioinformatically predicted to encode a portion of a natural protein.

As used herein, the term "peptide" refers to a sequence of amino acids. The sequences referred to as

PEPTIDE SEQ ID NOS.: are the predicted peptide sequences that would be translated from one of the exons, or a portion thereof set out in exon SEQ ID NOS.:. The codons encoding the peptide are wholly contained within the exon.

As used herein, a "portions" of a defined nucleotide sequence or sequences can be and, preferably, are fragments unique to that sequence or to one or a combination of those sequences. A fragment unique to a nucleic acid molecule is one that is a signature for the 10 larger nucleic acid molecule.

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As used herein, the phrase "expression of a probe" and its linguistic variants means that the ORF present within the probe, or its complement, is present within a target mRNA.

As used herein, "stringent conditions" refers to parameters well known to those skilled in the art. When a nucleic acid molecule is said to be hybridisable to another of a given sequence under "stringent conditions" it is meant that it is homologous to the given sequence.

As used herein, the phrase "specific binding 20 pair" intends a pair of molecules that bind to one another with high specificity. Binding pairs are said to exhibit specific binding when they exhibit avidity of at least 107, preferably at least 108, more preferably at least 109 liters/mole. Nonlimiting examples of specific binding pairs are: antibody and antigen; biotin and avidin; and biotin and streptavidin.

As used herein with respect to the visual display of annotated genomic sequence, the term "rectangle" means 30 any geometric shape that has at least a first and a second border, wherein the first and second borders each are capable of mapping uniquely to a point of another visual object of the display.

As used herein, a "Mondrian" means a visual display in which a single genomic sequence is annotated

with predicted and experimentally confirmed functional information.

5 Brief Description of the Drawings

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The present invention is further illustrated with reference to the following non-limiting figures and examples in which:

10 FIG. 1 illustrates a process for predicting functional regions from genomic sequence, confirming the functional activity of such regions experimentally, and associating and displaying the data so obtained in meaningful and useful relationship to the original sequence data;

FIG. 2 further elaborates that portion of the process schematized in FIG. 1 for predicting functional regions from genomic sequence;

FIG. 3 illustrates a Mondrian visual display;
FIG. 4 presents a Mondrian showing a hypothetical annotated genomic sequence;

FIG. 5 is a histogram showing the distribution of ORF length and PCR products as obtained, with ORF length shown in black and PCR product length shown in dotted lines;

FIG. 6 is a histogram showing the distribution, among exons predicted according to the methods described, of expression as measured using simultaneous two color hybridization to a genome-derived single exon microarray.

The graph shows the number of sequence-verified products that were either not expressed ("0"), expressed in one or more but not all tested tissues ("1" - "9"), or expressed in all tissues tested ("10");

FIG. 7 is a pictorial representation of the expression of verified sequences that showed expression

with signal intensity greater than 3 in at least one tissue, with: FIG. 7A showing the expression as measured by microarray hybridization in each of the 10 measured tissues, and the expression as measured "bioinformatically" by query of EST, NR and SwissProt databases; with FIG. 7B showing the legend for display of physical expression (ratio) in FIG. 7A; and with FIG. 7C showing the legend for scoring EST hits as depicted in FIG. 7A;

FIG. 8 shows a comparison of normalized CY3

10 signal intensity for arrayed sequences that were identical to sequences in existing EST, NR and SwissProt databases or that were dissimilar (unknown), where black denotes the signal intensity for all sequence-verified products with a BLAST Expect ("E") value of greater than 1e-30 (1 x 10⁻³⁰)

15 ("unknown") and a dotted line denotes sequence-verified spots with a BLAST expect ("E") value of less than 1e-30 (1 x 10⁻³⁰) ("known");

FIG. 9 presents a Mondrian of BAC AC008172 (bases 25,000 to 130,000), containing the carbamyl phosphate 20 synthetase gene (AF154830.1); and

FIG. 10 is a Mondrian of BAC A049839.

Methods and Apparatus for Predicting, Confirming,
 Annotating, and Displaying Functional Regions From Genomic
 Sequence Data

FIG. 1 is a flow chart illustrating in broad outline a process for predicting functional regions from genomic sequence, confirming and characterizing the functional activity of such regions experimentally, and then associating and displaying the information so obtained in meaningful and useful relationship to the original sequence data.

The initial input into process 10 of the present

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invention is drawn from one or more databases 100 containing genomic sequence data. Because genomic sequence is usually obtained from subgenomic fragments, the sequence data typically will be stored in a series of records corresponding to these subgenomic sequenced fragments. Some fragments will have been catenated to form larger contiguous sequences ("contigs"); others will not. A finite percentage of sequence data in the database will typically be erroneous, consisting inter alia of vector sequence, sequence created from aberrant cloning events, sequence of artificial polylinkers, and sequence that was erroneously read.

Each sequence record in database 100 will minimally contain as annotation a unique sequence

identifier (accession number), and will typically be annotated further to identify the date of accession, species of origin, and depositor. Because database 100 can contain nongenomic sequence, each sequence will typically be annotated further to permit query for genomic sequence.

Chromosomal origin, optionally with map location, can also be present. Data can be, and over time increasingly will be, further annotated with additional information, in part through use of the present invention, as described below. Annotation can be present within the data records, in information external to database 100 and linked to the records thereto, or through a combination of the two.

Databases useful as genomic sequence database 100 in the present invention include GenBank, and particularly include several divisions thereof, including the

30 htgs(draft), NT (nucleotide, command line), and NR (nonredundant) divisions. GenBank is produced by the National Institutes of Health and is maintained by the National Center for Biotechnology Information (NCBI).

Databases of genomic sequence from species other than

35 human, such as mouse, rat, Arabidopsis, C. elegans, C.

brigsii, Drosophila, zebra fish, and other higher eukaryotic organisms will also prove useful as genomic sequence database 100.

Genomic sequence obtained by query of genomic

5 sequence database 100 is then input into one or more
processes 200 for identification of regions therein that
are predicted to have a biological function as specified by
the user. Such functions include, but are not limited to,
encoding protein, regulating transcription, regulating

10 message transport after transcription into mRNA, regulating
message splicing after transcription into mRNA, of
regulating message degradation after transcription into
mRNA, and the like. Other functions include directing
somatic recombination events, contributing to chromosomal

15 stability or movement, contributing to allelic exclusion or
X chromosome inactivation, and the like.

The particular genomic sequence to be input into process 200 will depend upon the function for which relevant sequence is to be identified as well as upon the approach chosen for such identification. Process step 200 can be iterated to identify different functions within a given genomic region. In such case, the input often will be different for the several iterations.

Sequences predicted to have the requisite

25 function by process 200 are then input into process 300,
where a subset of the input sequences suitable for
experimental confirmation is identified. Experimental
confirmation can involve physical and/or bioinformatic
assay. Where the subsequent experimental assay is
30 bioinformatic, rather than physical, there are fewer
constraints on the sequences that can be tested, and in
this latter case therefore process 300 can output the
entirety of the input sequence.

The subset of sequences output from process 300 is then used in process 400 for experimental verification

and characterization of the function predicted in process 200, which experimental verification can, and often will, include both physical and bioinformatic assay.

Process 500 annotates the sequence data with the functional information obtained in the physical and/or bioinformatic assays of process 400. Such annotation can be done using any technique that usefully relates the functional information to the sequence, as, for example, by incorporating the functional data into the sequence data record itself, by linking records in a hierarchical or relational database, by linking to external databases, by a combination thereof, or by other means well known within the database arts. The data can even be submitted for incorporation into databases maintained by others, such as GenBank, which is maintained by NCBI.

As further noted in FIG. 1, additional annotation can be input into process 500 from external sources 600.

The annotated data is then displayed in process 800, either before, concomitantly with, or after optional storage 700 on nontransient media, such as magnetic disk, optical disc, magnetooptical disk, flash memory, or the like.

FIG. 1 shows that the experimental data output from process 400 can be used in each preceding step of process 10: e.g., facilitating identification of functional sequences in process 200, facilitating identification of an experimentally suitable subset thereof in process 300, and facilitating creation of physical and/or informational substrates for, and performance of subsequent assay, of functional sequences in process 400.

Information from each step can be passed directly to the succeeding process, or stored in permanent or interim form prior to passage to the succeeding process.

Often, data will be stored after each, or at least a plurality, of such process steps. Any or all process steps

can be automated.

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FIG. 2 further elaborates the prediction of functional sequence within genomic sequence according to process 200.

5 Genomic sequence database 100 is first queried 20 for genomic sequence.

The sequence required to be returned by query 20 will depend, in the first instance, upon the function to be identified.

For example, genomic sequences that function to encode protein can be identified inter alia using gene prediction approaches, comparative sequence analysis approaches, or combinations of the two. In gene prediction analysis, sequence from one genome is input into process 200 where at least one, preferably a plurality, of algorithmic methods are applied to identify putative coding regions. In comparative sequence analysis, by contrast, corresponding, e.g., syntenic, sequence from a plurality of sources, typically a plurality of species, is input into process 200, where at least one, possibly a plurality, of algorithmic methods are applied to compare the sequences and identify regions of least variability.

The exact content of query 20 will also depend upon the database queried. For example, if the database contains both genomic and nongenomic sequence, perhaps derived from multiple species, and the function to be determined is protein coding regions in human genomic sequence, the query will accordingly require that the sequence returned be genomic and derived from humans.

Query 20 can also incorporate criteria that compel return of sequence that meets operative requirements of the subsequent analytical method. Alternatively, or in addition, such operative criteria can be enforced in subsequent preprocess step 24.

For example, if the function sought to be

identified is protein coding, query 20 can incorporate criteria that return from genomic sequence database 100 only those sequences present within contigs sufficiently long as to have obviated substantial fragmentation of any given exon among a plurality of separate sequence fragments.

Such criteria can, for example, consist of a required minimal individual genomic sequence fragment length, such as 10 kb, more typically 20 kb, 30 kb, 40kb, and preferably 50 kb or more, as well as an optional further or alternative requirement that sequence from any given clone, such as a bacterial artificial chromosome ("BAC"), be presented in no more than a finite maximal number of fragments, such as no more than 20 separate pieces, more typically no more than about 10 - 12 fragments.

Results using the present invention have shown that genomic sequence from bacterial artificial chromosomes (BACs) is sufficient for gene prediction analysis according to the present invention if the sequence is at least 50 kb in length, and if additionally the sequence from any given BAC is presented in fewer than 15, and preferably fewer than 10, fragments. Accordingly, query 20 can incorporate a requirement that data accessioned from BAC sequencing be in fewer than 15, preferably fewer than 10, fragments.

An additional criterion that can be incorporated into the query can be the date, or range of dates, of sequence accession. Although the process has been described above as if genomic sequence database 100 were static, it is of course understood that the genomic sequence databases need not be static, and indeed are typically updated on a frequent, even hourly, basis. Thus, as further described in Examples 1 and 2, infra, it is possible to query the database for newly added sequence, either newly added after an absolute date, or newly added

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relative to a prior analysis performed using the methods and apparatus of the present invention. In this way, the process herein described can incorporate a dynamic, temporal component.

One utility of such temporal limitation is to 5 identify, from newly accessioned genomic sequence, the presence of novel genes, particularly those not previously identified by EST sequencing (or other sequencing efforts that are similarly based upon gene expression). As further 10 described in Example 1, such an approach has shown that newly accessioned human genomic sequence, when analyzed for sequences that function to encode protein, readily identifies genes that are novel over those in existing EST and other expression databases. This makes the methods of 15 the present invention extremely powerful gene discovery tools. And as would be appreciated, such gene discovery can be performed using genomic sequence from species other than human.

If query 20 incorporates multiple criteria, such 20 as above-described, the multiple criteria can be performed as a series of separate queries or as a single query, depending in part upon the query language, the complexity of the query, and other considerations well known in the database arts.

If query 20 returns no genomic sequence meeting the query criteria, the negative result can be reported by process 22, and process 200 (and indeed, entire process 10) ended 23, as shown. Alternatively, or in addition to report and termination of the initial inquiry, a new query 30 20 can be generated that takes into account the initial negative result.

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When query 20 returns sequence meeting the query criteria, the returned sequence is then passed to optional preprocessing 24, suitable and specific for the desired 35 analytical approach and the particular analytical methods

thereof to be used in process 25.

Preprocessing 24 can include processes suitable for many approaches and methods thereof, as well as processes specifically suited for the intended subsequent analysis.

Preprocessing 24 suitable for most approaches and methods will include elimination of sequence irrelevant to, or that would interfere with, the subsequent analysis.

Such sequence includes repetitive sequence, such as Alu

repeats and LINE elements, vector sequence, artificial sequence, such as artificial polylinkers, and the like.

Such removal can readily be performed by identification and subsequent masking of the undesired sequence.

Identification can be effected by comparing the
genomic sequence returned by query 20 with public or
private databases containing known repetitive sequence,
vector sequence, artificial sequence, and other artifactual
sequence. Such comparison can readily be done using
programs well known in the art, such as CROSS_MATCH, or by
proprietary sequence comparison programs the engineering of
which is well within the skill in the art.

Alternatively, or in addition, undesirable, including artifactual, sequence can be identified algorithmically without comparison to external databases

25 and thereafter removed. For example, synthetic polylinker sequence can be identified by an algorithm that identifies a significantly higher than average density of known restriction sites. As another example, vector sequence can be identified by algorithms that identify nucleotide or codon usage at variance with that of the bulk of the genomic sequence.

Once identified, undesired sequence can be removed. Removal can usefully be done by masking the undesired sequence as, for example, by converting the specific nucleotide references to one that is unrecognized

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by the subsequent bioinformatic algorithms, such as "X". Alternatively, but at present less preferred, the undesired sequence can be excised from the returned genomic sequence, leaving gaps.

Preprocessing 24 can further include selection from among duplicative sequences of that one sequence of highest quality. Higher quality can be measured as a lower percentage of, fewest number of, or least densely clustered occurrence of ambiguous nucleotides, defined as those 10 nucleotides that are identified in the genomic sequence using symbols indicating ambiguity. Higher quality can also or alternatively be valued by presence in the longest contig.

Preprocessing 24 can, and often will, also 15 include formatting of the data as specifically appropriate for passage to the analytical algorithms of process 25. Such formatting can and typically will include, inter alia, addition of a unique sequence identifier, either derived from the original accession number in genomic sequence 20 database 100, or newly applied, and can further include additional annotation. Formatting can include conversion from one to another sequence listing standard, such as 🐭 conversion to or from FASTA or the like, depending upon the input expected by the subsequent process.

Preprocessing, which can be optional depending upon the function desired to be identified and the informational requirements of the methods for effecting such identification, is followed by sequence processing 25, where sequences with the desired function are identified 30 within the genomic sequence.

As mentioned above, such functions can include, but are not limited to, encoding protein, regulating transcription, regulating message transport after transcription into mRNA, regulating message splicing after 35 transcription, of regulating message degradation, and the

like. Other functions include directing somatic recombination events, contributing to chromosomal stability or movement, contributing to allelic exclusion or X chromosome inactivation, or the like.

5 The methods of the present invention are particularly useful for gene discovery, that is, for identifying, from genomic sequence, regions that function to encode genes, and in a particularly useful embodiment, for identifying regions that function to encode genes not 10 hitherto identified by expression-based or directed cloning and sequencing. In conjunction with verification using the novel single exon microarrays of the present invention, as further described below, the methods herein described become powerful gene discovery tools.

Accordingly, in a preferred embodiment of the present invention, process 25 is used to identify putative coding regions. Two preferred approaches in process 25 for identifying sequence that encodes putative genes are gene prediction and comparative sequence analysis.

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Gene prediction can be performed using any of a number of algorithmic methods, embodied in one or more software programs, that identify open reading frames (ORFs) using a variety of heuristics, such as GRAIL, DICTION, and GENEFINDER. Comparative sequence analysis similarly can be 25 performed using any of a variety of known programs that identify regions with lower sequence variability.

As further described in Example 1, below, gene finding software programs yield a range of results. For the newly accessioned human genomic sequence input in 30 Example 1, for example, GRAIL identified the greatest percentage of genomic sequence as putative coding region, 2% of the data analyzed; GENEFINDER was second, calling 1%; and DICTION yielded the least putative coding region, with 0.8% of genomic sequence called as coding region.

35 Increased reliability can be obtained when

consensus is required among several such methods. Although discussed herein particularly with respect to exon calling, consensus among methods will in general increase reliability of predicting other functions as well.

Thus, as indicated by query 26, sequence processing 25, optionally with preprocessing 24, can be repeated with a different method, with consensus among such iterations determined and reported in process 27.

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Process 27 compares the several outputs for a given input genomic sequence and identifies consensus among the separately reported results. The consensus itself, as well as the sequence meeting that consensus, is then stored in process 29a, displayed in process 29b, and/or output to process 300 for subsequent identification of a subset thereof suitable for assay.

Multiple levels of consensus can be calculated and reported by process 27. For example, as further described in Example 1, infra, process 27 can report consensus as between all specific pairs of methods of gene prediction, as consensus among any one or more of the pairs of methods of gene prediction, or as among all of the gene prediction algorithms used. Thus, in Example 1, process 27 reported that GRAIL and GENEFINDER programs agreed on 0.7% of genomic sequence, that GRAIL and DICTION agreed on 0.5% of genomic sequence, and that the three programs together agreed on 0.25% of the data analyzed. Put another way, 0.25% of the genomic sequence was identified by all three of the programs as containing putative coding region.

Furthermore, consensus can be required among different approaches to identifying a chosen function.

For example, if the function desired to be identified is coding of protein sequence, and a first used approach to exon calling is gene prediction, the process can be repeated on the same input sequence, or subset thereof, with another approach, such as comparative

sequence analysis. In such a case, where comparative sequence analysis follows gene prediction, the comparison can be performed not only on genomic nucleic acid sequence, but additionally or alternatively can be performed on the 5 predicted amino acid sequence translated from the ORFs prior identified by the gene prediction approach.

Although shown as an iterative process, the multiple analyses required to achieve consensus can be done in series, in parallel, or some combination thereof.

Predicted functional sequence, optionally representing a consensus among a plurality of methods and approaches for determination thereof, is passed to process 300 for identification of a subset thereof for functional assay.

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In the preferred embodiment of the methods of the present invention, wherein the function sought to be identified is protein coding, process 300 is used to identify a subset thereof suitable for experimental verification by physical and/or bioinformatic approaches.

For example, putative ORFs identified in process . 200 can be classified, or binned, bioinformatically into putative genes. This binning can be based inter alia upon consideration of the average number of exons/gene in the species chosen for analysis, upon density of exons that 25 have been called on the genomic sequence, and other empirical rules. Thereafter, one or more among the genespecific ORFs can be chosen for subsequent use in gene expression assay.

Where such subsequent gene expression assay uses 30 amplified nucleic acid, considerations such as desired amplicon length, primer synthesis requirements, putative exon length, sequence GC content, existence of possible secondary structure, and the like can be used to identify and select those ORFs that appear most likely successfully 35 to amplify. Where subsequent gene expression assay relies

upon nucleic acid hybridization, whether or not using amplified product, further considerations involving hybridization stringency can be applied to identify that subset of sequences that will most readily permit sequences specific discrimination at a chosen hybridization and wash stringency. One particular such consideration is avoidance of putative exons that span repetitive sequence; such sequence can hybridize spuriously to nonspecific message, reducing specific signal in the hybridization.

For bioinformatic assay, there are fewer constraints on the sequences that can be tested experimentally, and in this latter case therefore process 300 can output the entirety of the input sequence.

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The subset of sequences identified by process 300 as suitable for use in assay is then used in process 400 to create the physical and/or informational substrate for experimental verification of the predictions made in process 200, and thereafter to assay those substrates.

As mentioned, the methods of the present
invention are particularly useful for identifying potential
coding regions within genomic sequence. In a preferred
embodiment of process 400, therefore, the expression of the
sequences predicted to encode protein is verified. The
combination of the predictive and experimental methods
provides a powerful gene discovery engine.

Thus, in another aspect, the present invention provides methods and apparatus for verifying the expression of putative genes identified within genomic sequence. In particular, the invention provides a novel method of verifying gene expression in which expression of predicted ORFs is measured and confirmed using a novel type of nucleic acid microarray, the genome-derived single exon nucleic acid microarrays of the present invention.

Putative ORFs as predicted by a consensus of gene calling, particularly gene prediction, algorithms in

process 200, and as further identified as suitable by process 300, are amplified from genomic DNA using the polymerase chain reaction (PCR). Although PCR is conveniently used, other amplification approaches can also be used.

Amplification schemes can be designed to capture the entirety of each predicted ORF in an amplicon with minimal additional (that is, intronic or intergenic) sequence. Because ORFs predicted from human genomic sequence using the methods of the present invention differ in length, such an approach results in amplicons of varying length.

However, most predicted ORFs are shorter than 500 bp in length, and although amplicons of at least about 100 or 200 base pairs can be immobilized as probes on nucleic acid microarrays, early experimental results using the methods of the present invention have suggested that longer amplicons, at least about 400 or 500 base pairs, are more effective. Furthermore, certain advantages derive from application to the microarray of amplicons of defined size.

Therefore, amplification schemes can alternatively, and preferably, be designed to amplify regions of defined size, preferably at least about 300, 400 or 500 bp, centered about each predicted ORF. Such an approach results in a population of amplicons of limited size diversity, but that typically contain intronic and/or intergenic nucleic acid in addition to putative ORF.

Conversely, somewhat fewer than 10% of ORFs predicted from human genomic sequence according to the methods of the present invention exceed 500 bp in length. Portions of such extended ORFs, preferably at least about 300,400 or 500 bp in length, can be amplified. However, it has been discovered that the percentage success at amplifying pieces of such ORFs is low, and that such putative exons are more effectively amplified when larger

fragments, at least about 1000 or 1500 bp, and even as large as 2000 bp are amplified.

The putative ORFs selected in process 300 are thus input into one or more primer design programs, such as PRIMER3 (available online for use at http://www-genome.wi.mit.edu/cgi-bin/primer/), with a goal of amplifying at least about 500 base pairs of genomic sequence centered within or about ORFs predicted to be no more than about 500 bp, or at least about 1000 - 1500 bp of genomic sequence for ORFs predicted to exceed 500 bp in length, and the primers synthesized by standard techniques. Primers with the requisite sequences can be purchased commercially or synthesized by standard techniques.

Conveniently, a first predetermined sequence can

be added commonly to the ORF-specific 5' primer and a

second, typically different, predetermined sequence

commonly added to each 3' ORF-unique primer. This serves

to immortalize the amplicon, that is, serves to permit

further amplification of any amplicon using a single set of

primers complementary respectively to the common 5' and

common 3' sequence elements. The presence of these

"universal" priming sequences further facilitates later

sequence verification, providing a sequence common to all

amplicons at which to prime sequencing reactions. The

common 5' and 3' sequences further serve to add a cloning

site should any of the ORFs warrant further study.

Such predetermined sequence is usefully at least about 10, 12 or 15 nt in length, and usually does not exceed about 25 nt in length. The "universal" priming sequences used in the examples presented *infra* were each 16 nt long.

The genomic DNA to be used as substrate for amplification will come from the eukaryotic species from which the genomic sequence data had originally been obtained, or a closely related species, and can

conveniently be prepared by well known techniques from somatic or germline tissue or cultured cells of the organism. See, e.g., Short Protocols in Molecular Biology: A Compendium of Methods from Current Protocols in

Molecular Biology, Ausubel et al. (eds.), 4th edition (April 1999), John Wiley & Sons (ISBN: 047132938X) and Maniatis et al., Molecular Cloning: A Laboratory Manual, 2nd edition (December 1989), Cold Spring Harbor Laboratory Press (ISBN: 0879693096). Many such prepared genomic DNAs are available commercially, with the human genomic DNAs additionally having certification of donor informed consent.

Although the intronic and intergenic material flanking putative coding regions in the amplicons could potentially interfere with hybridizations during microarray experiments, we have found, surprisingly, that differential expression ratios are not significantly affected. Rather, the predominant effect of exon size is to alter the absolute signal intensity, rather than its ratio. Equally surprising, the art had suggested that single exon probes would not provide sufficient signal intensity for high stringency hybridization analyses; we find that such probes not only provide adequate signal, but have substantial advantages, as herein described.

25 After partial purification, as by size exclusion spin column, with or without confirmation as to amplicon quality as by gel electrophoresis, each amplicon (single exon probe) is disposed in an array upon a support substrate.

Methods for creating microarrays by deposition and fixation of nucleic acids onto support substrates are well known in the art (Reviewed by Schena et al., see above).

Typically, the support substrate will be glass, although other materials, such as amorphous or crystalline

silicon or plastics. Such plastics include
polymethylacrylic, polyethylene, polypropylene,
polyacrylate, polymethylmethacrylate, polyvinylchloride,
polytetrafluoroethylene, polystyrene, polycarbonate,

5 polyacetal, polysulfone, celluloseacetate,
cellulosenitrate, nitrocellulose, or mixtures thereof, can
also be used. Typically, the support will be rectangular,
although other shapes, particularly circular disks and even
spheres, present certain advantages. Particularly
10 advantageous alternatives to glass slides as support
substrates for array of nucleic acids are optical discs, as
described in WO 98/12559.

The amplified nucleic acids can be attached covalently to a surface of the support substrate or, more typically, applied to a derivatized surface in a chaotropic agent that facilitates denaturation and adherence by presumed noncovalent interactions, or some combination thereof.

Robotic spotting devices useful for arraying

nucleic acids on support substrates can be constructed
using public domain specifications (The MGuide, version
2.0, http://cmgm.stanford.edu/pbrown/mguide/index.html), or
can conveniently be purchased from commercial sources
(MicroArray GenII Spotter and MicroArray GenIII Spotter,

Molecular Dynamics, Inc., Sunnyvale, CA). Spotting can
also be effected by printing methods, including those using
ink jet technology.

As is well known in the art, microarrays typically also contain immobilized control nucleic acids.

For controls useful in providing measurements of background signal for the genome-derived single exon microarrays of the present invention, a plurality of *E. coli* genes can readily be used. As further described in Example 1, 16 or 32 *E. coli* genes suffice to provide a robust measure of background noise in such microarrays.

As is well known in the art, the amplified product disposed in arrays on a support substrate to create a nucleic acid microarray can consist entirely of natural nucleotides linked by phosphodiester bonds, or

5 alternatively can include either nonnative nucleotides, alternative internucleotide linkages, or both, so long as complementary binding can be obtained in the hybridization. If enzymatic amplification is used to produce the immobilized probes, the amplifying enzyme will impose certain further constraints upon the types of nucleic acid analogs that can be generated.

Although particularly described herein as using high density microarrays constructed on planar substrates, the methods of the present invention for confirming the expression of ORFs predicted from genomic sequence can use any of the known types of microarrays, as herein defined, including lower density planar arrays, and microarrays on nonplanar, nonunitary, distributed substrates.

using hybridization to lower density arrays, such as those constructed on membranes, such as nitrocellulose, nylon, and positively-charged derivatized nylon membranes. Further, gene expression can also be confirmed using nonplanar, bead-based microarrays such as are described in Brenner et al., Proc. Natl. Acad. Sci. USA 97(4):166501670 (2000); U.S. Patent No. 6,057,107; and U.S. Patent No. 5,736,330. In theory, a packed collection of such beads provides in aggregate a higher density of nucleic acid probe than can be achieved with spotting or lithography techniques on a single planar substrate.

Planar microarrays on solid substrates, however, provide certain useful advantages, including high throughput and compatibility with existing readers. For example, each standard microscope slide can include at least 1000, typically at least 2000, preferably 5000 and

upto 10,000 - 50,000 or more nucleic acid probes of discrete sequence. The number of sequences deposited will depend on their required application.

Each putative gene can be represented in the

array by a single predicted ORF. Alternatively, genes can
be represented by more than one predicted ORF. For
purposes of measuring differential splicing, more than one
predicted ORF will be provided for a putative gene. And as
is well known in the art, each probe of defined sequence,
representing a single predicted ORF, can be deposited in a
plurality of locations on a single microarray to provide
redundancy of signal.

The genome-derived single exon microarrays described above differ in several fundamental and

15 advantageous ways from microarrays presently used in the gene expression art, including (1) those created by deposition of mRNA-derived nucleic acids, (2) those created by in situ synthesis of oligonucleotide probes, and (3) those constructed from yeast genomic DNA.

Most nucleic acid microarrays that are in use for study of eukaryotic gene expression have as immobilized probes nucleic acids that are derived — either directly or indirectly — from expressed message. As discussed above, it is common, for example, for such microarrays to be derived from cDNA/EST libraries, either from those previously described in the literature, see Lennon et al., or from the de novo construction of "problem specific" libraries targeted at a particular biological question, R.S. Thomas et al., Cancer Res. (in press). Such microarrays are herein collectively denominated "EST microarrays".

Such EST microarrays by definition can measure expression only of those genes found in EST libraries, shown herein to represent only a fraction of expressed genes. Furthermore, such libraries — and thus microarrays

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based thereupon — are biased by the tissue or cell type of message origin, by the expression levels of the respective genes within the tissues, and by the ability of the message successfully to have been reverse-transcribed and cloned.

Thus, as further discussed in Example 1, the methods of the present invention enable sequences that do not appear in EST or other expression databases to be determined - subsequently arrayed for expression measurements could not, therefore, have been represented as probes on an EST microarray. And as further demonstrated in the examples, infra, the remaining population of genes identified from genomic sequence by the methods of the present invention - that is, the one third of sequences that had previously been accessioned in EST or other expression databases - are biased toward genes with higher expression levels.

Representation of a message in an EST and/or cDNA library depends upon the successful reverse transcription, optionally but typically with subsequent successful cloning, of the message. This introduces substantial bias into the population of probes available for arraying in EST microarrays.

In contrast, neither reverse transcription nor cloning is required to produce the probes arrayed on the genome-derived single exon microarrays of the present invention. And although the ultimate deposition of a probe on the genome-derived single exon microarray of the present invention depends upon a successful amplification from genomic material, a priori knowledge of the sequence of the desired amplicon affords greater opportunity to recover any given probe sequence recalcitrant to amplification than is afforded by the requirement for successful reverse transcription and cloning of unknown message in EST approaches.

Thus, the genome-derived single exon microarrays

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of the present invention present a far greater diversity of probes for measuring gene expression, with far less bias, than do EST microarrays presently used in the art.

As a further consequence of their ultimate origin 5 from expressed message, the probes in EST microarrays often contain poly-A (or complementary poly-T) stretches derived from the poly-A tail of mature mRNA. These homopolymeric stretches contribute to cross-hybridization, that is, to a spurious signal occasioned by hybridization to the 10 homopolymeric tail of a labeled cDNA that lacks sequence homology to the gene-specific portion of the probe.

In contrast, the probes arrayed in the genomederived single exon microarrays of the present invention lack homopolymeric stretches derived from message 15 polyadenylation, and thus can provide more specific signal. Typically, at least about 50, 60 or 75% of the probes on the genome-derived single exon microarrays of the present invention lack homopolymeric regions consisting of A or T, where a homopolymeric region is defined for purposes herein 20 as stretches of 25 or more, typically 30 or more, identical nucleotides.

A further distinction, which also affects the specificity of hybridization, is occasioned by the typical derivation of EST microarray probes from cloned material. 25 Because much of the probe material disposed as probes on EST microarrays is excised or amplified from plasmid, phage, or phagemid vectors, EST microarrays typically include a fair amount of vector sequence, more so when the probes are amplified, rather than excised, from the vector.

In contrast, the vast majority of probes in the genome-derived single exon microarrays of the present invention contain no prokaryotic or bacteriophage vector sequence, having been amplified directly or indirectly from genomic DNA. Typically, therefore, at least about 50, 60, 35 70 or 80% or more of individual exon-including probes

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disposed on a genome-derived single exon microarray of the present invention lack vector sequence, and particularly lack sequences drawn from plasmids and bacteriophage. Preferably, at least about 85, 90 or more than 90% of exonincluding probes in the genome-derived single exon microarray of the present invention lack vector sequence. With attention to removal of vector sequences through preprocessing 24, percentages of vector-free exon-including probes can be as high as 95 - 99%. The substantial absence of vector sequence from the genome-derived single exon microarrays of the present invention results in greater specificity during hybridization, since spurious cross-hybridization to a probe vector sequence is reduced.

As a further consequence of excision or

15 amplification of probes from vectors in construction of EST
microarrays, the probes arrayed thereon often contain
artificial sequence, derived from vector polylinker
multiple cloning sites, at both 5' and 3' ends. The probes
disposed upon the genome-derived single exon microarrays

20 need have no such artificial sequence appended thereto.

As mentioned above, however, the ORF-specific primers used to amplify putative ORFs can include artificial sequences, typically 5' to the ORF-specific primer sequence, useful for "universal" (that is, independent of ORF sequence) priming of subsequent amplification or sequencing reactions. When such "universal" 5' and/or 3' priming sequences are appended to the amplification primers, the probes disposed upon the genome-derived single exon microarray will include artificial sequence similar to that found in EST microarrays. However, the genome-derived single exon microarray of the present invention can be made without such sequences, and if so constructed, presents an even smaller amount of nonspecific sequence that would contribute to nonspecific hybridization.

Yet another consequence of typical use of cloned material as probes in EST microarrays is that such microarrays contain probes that result from cloning artifacts, such as chimeric molecules containing coding region of two separate genes. Derived from genomic material, typically not thereafter cloned, the probes of the genome-derived single exon microarrays of the present invention lack such cloning artifacts, and thus provide greater specificity of signal in gene expression measurements.

A further consequence of the cloned origin of probes on many EST microarrays is that the individual probes often have disparate sizes, which can cause the optimal hybridization stringency to vary among probes on a single microarray. In contrast, as discussed above, the probes arrayed on the genome-derived single exon microarrays of the present invention can readily be designed to have a narrow distribution in sizes, with the range of probe sizes no greater than about 10% of the average size, typically no greater than about 5% of the average probe size.

Because of their origin from fully- or partiallyspliced message, probes disposed upon EST arrays will often
include multiple exons. The percentage of such exon25 spanning probes in an EST microarray can be calculated, on
average, based upon the predicted number of exons/gene for
the given species and the average length of the immobilized
probes. For human genes, the near-complete sequence of
human chromosome 22, Dunham et al., Nature 402(6761):489-95
30 (1999), predicts that human genes average 5.5 exons/gene.
Even with probes of 200 - 500 bp, the vast majority of
human EST microarray probes include more than one exon.

In contrast, by virtue of their origin from algorithmically identified ORFs in genomic sequence, the probes in the genome-derived single exon microarrays of the

present invention can consist of individual exons. Thus, in contrast to EST microarrays, at least about 50, 60, 70, 75, 80, 85, 95 or 99% of probes deposited in the genomederived microarray of the present invention consist of, or include, no more than one predicted ORF.

This provides the ability, not readily achieved using EST microarrays, to use the genome-derived single exon microarrays of the present invention to measure tissue-specific expression of individual exons, which in turn allows differential splicing events to be detected and characterized, and in particular, allows the correlation of differential splicing to tissue-specific expression patterns.

Furthermore, the exons that are represented in

15 EST microarrays are often biased toward the 3' or 5' end of
their respective genes, since sequencing strategies used
for EST identification are so biased. In contrast, no such
3' or 5' bias necessarily inheres in the selection of exons
for disposition on the genome-derived single exon

20 microarrays of the present invention.

Conversely, the probes provided on the genomederived single exon microarrays of the present invention
typically, but need not necessarily, include intronic
and/or intergenic sequence that is absent from EST

25 microarrays, which are derived from mature mRNA.
Typically, at least about 50, 60, 70, 80 or 90% of the
exon-including probes on the genome-derived single exon
microarrays of the present invention include sequence drawn
from noncoding regions. As discussed above, the additional
30 presence of noncoding region does not significantly
interfere with measurement of gene expression, and provides
the additional opportunity to assay prespliced RNA, and
thus measure such phenomena such as nuclear export control.

The genome-derived single exon microarrays of the present invention are also quite different from in situ

synthesis microarrays, where probe size is severely constrained by inadequacies in the photolithographic synthesis process.

Typically, probes arrayed on in situ synthesis

5 microarrays are limited to a maximum of about 25 bp. As a
well known consequence, hybridization to such chips must be
performed at low stringency. In order, therefore, to
achieve unambiguous sequence-specific hybridization
results, the in situ synthesis microarray requires

10 substantial redundancy, with concomitant programmed
arraying for each probe of probe analogues with altered
(i.e., mismatched) sequence.

In contrast, the longer probe length of the genome-derived single exon microarrays of the present

15 invention allows much higher stringency hybridization and wash. Typically, therefore, exon-including probes on the genome-derived single exon microarrays of the present invention average at least about 100, 200, 300, 400 or 500 bp in length. By obviating the need for substantial probe redundancy, this approach permits a higher density of probes for discrete exons or genes to be arrayed on the microarrays of the present invention than can be achieved for in situ synthesis microarrays.

A further distinction is that the probes in in

25 situ synthesis microarrays typically are covalently linked
to the substrate surface. In contrast, the probes disposed
on the genome-derived microarray of the present invention
typically are, but need not necessarily be, bound
noncovalently to the substrate.

Furthermore, the short probe size on in situ microarrays causes large percentage differences in the melting temperature of probes hybridized to their complementary target sequence, and thus causes large percentage differences in the theoretically optimum stringency across the array as a whole.

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In contrast, the larger probe size in the microarrays of the present invention create lower percentage differences in melting temperature across the range of arrayed probes.

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A further significant advantage of the microarrays of the present invention over in situ synthesized arrays is that the quality of each individual probe can be confirmed before deposition. In contrast, the quality of probes cannot be assessed on a probe-by-probe 10 basis for the *in situ* synthesized microarrays presently being used.

The genome-derived single exon microarrays of the present invention are also distinguished over, and present substantial benefits over, the genome-derived microarrays 15 from lower eukaryotes such as yeast. Lashkari et al., Proc. Natl. Acad. Sci. USA 94:13057-13062 (1997).

Only about 220 - 250 of the 6100 or so nuclear genes in Saccharomyces cerevisiae - that is, only about 4 - 5% - have standard, spliceosomal, introns, Lopez et al., 20 Nucl. Acids Res. 28:85-86 (2000); Spingola et al., RNA 5(2):221-34 (1999). Furthermore, the entire yeast genome has already been sequenced. These two facts permit the ready amplification and disposition of single-ORF amplicons on such microarray without the requirement for antecedent 25 use of gene prediction and/or comparative sequence analyses.

Thus, a significant aspect of the present invention is the ability to identify and to confirm expression of predicted coding regions in genomic sequence 30 drawn from eukaryotic organisms that have a higher percentage of genes having introns than do yeast such as Saccharomyces cerevisiae, particularly in genomic sequence drawn from eukaryotes in which at least about 10, 20 or 50% of protein-encoding genes have introns. In preferred 35 embodiments, the methods and apparatus of the present

invention are used to identify and confirm expression of novel genes from genomic sequence of eukaryotes in which the average number of introns per gene is at least about one, two or three or more.

After the physical substrate is prepared, experimental verification of predicted function is performed.

In a preferred embodiment of the present invention, where the function sought to be identified in genomic sequence is protein coding, experimental verification is performed by measuring expression of the putative ORFs, typically through nucleic acid hybridization experiments, and in particularly preferred embodiments, through hybridization to genome-derived single exon microarrays prepared as above- described.

Expression is conveniently measured and expressed for each probe in the microarray as a ratio of the expression measured concurrently in a plurality of mRNA sources, according to techniques well known in the

20 microarray art, Reviewed in Schena et al., and as further described in Example 2, below. The mRNA source for the reference against which specific expression is measured can be drawn from a homogeneous mRNA source, such as a single cultured cell-type, or alternatively can be heterogeneous,

25 as from a pool of mRNA derived from multiple tissues and/or cell types, as further described in Example 2, infra.

mRNA can be prepared by standard techniques, see
Ausubel et al. and Maniatis et al., or purchased
commercially. The mRNA is then typically reverse
transcribed in the presence of labeled nucleotides: the
index source (that in which expression is desired to be
measured) is reverse transcribed in the presence of
nucleotides labeled with a first label, typically a
fluorophore (fluorochrome; fluor; fluorescent dye); the
reference source is reverse transcribed in the presence of

a second label, typically a fluorophore, typically fluorometrically-distinguishable from the first label. As further described in Example 2, infra, Cy3 and Cy5 dyes prove particularly useful in these methods. After partial purification of the index and reference targets, hybridization to the probe array is conducted according to standard techniques, typically under a coverslip.

After wash, microarrays are conveniently scanned using a commercial microarray scanning device, such as a 10 Gen3 Scanner (Molecular Dynamics, Sunnyvale, CA). Data on expression is then passed, with or without interim storage, to process 500, where the results for each probe are related to the original sequence.

Often, hybridization of target material to the

genome-derived single exon microarray will identify certain
of the probes thereon as of particular interest. Thus, it
is often desirable that the user be able readily to obtain
sufficient quantities of an individual probe, either for
subsequent arrayed deposition upon an additional support
substrate, often as part of a microarray having a plurality
of probes so identified, or alternatively or additionally
as a solitary solid-phase or solution-phase probe, for
further use.

Thus, in another aspect, the present invention
25 provides compositions and kits for the ready production of
nucleic acids identical in sequence to, or substantially
identical in sequence to, probes on the genome-derived
single exon microarrays of the present invention.

In this aspect, a small quantity of each probe is 30 disposed, typically without attachment to substrate, in a spatially-addressable ordered set, typically one per well of a microtiter dish. Although a 96 well microtiter plate can be used, greater efficiency is obtained using higher density arrays, such as are provided by microtiter plates 35 having 384, 864, 1536, 3456, 6144, or 9600 wells, and

although microtiter plates having physical depressions (wells) are conveniently used, any device that permits addressable withdrawal of reagent from fluidlynoncommunicating areas can be used.

In this aspect of the invention, therefore, a fluidly noncommunicating addressable ordered set of individual probes, corresponding to those on a genomederived single exon microarray, is provided, with each probe in sufficient quantity to permit amplification, such 10 as by PCR. As earlier mentioned, the ORF-specific 5' primers used for genomic amplification can have a first common sequence added thereto, and the ORF-specific 3' primers used for genomic amplification can have a second, different, common sequence added thereto, thus permitting, 15 in this preferred embodiment, the use of a single set of 5' and 3' primers to amplify any one of the probes from the amplifiable ordered set.

Each discrete amplifiable probe can also be packaged with amplification primers, solutes, buffers, 20 etc., and can be provided in dry (e.g., lyophilized) form or wet, in the latter case typically with addition of agents that retard evaporation.

In another aspect of the present invention, a genome-derived single-exon microarray is packaged together 25 with such an ordered set of amplifiable probes corresponding to the probes, or one or more subsets of In alternative embodiments, the ordered probes, thereon. set of amplifiable probes is packaged separately from the genome-derived single exon microarray.

In some embodiments, the microarray and/or ordered probe set are further packaged with recordable media that provide probe identification and addressing information, and that can additionally contain annotation information, such as gene expression data. Such recordable 35 media can be packaged with the microarray, with the ordered

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probe set, or with both.

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If the microarray is constructed on a substrate that incorporates recordable media, such as is described in international patent application no. WO 98/12559, then 5 separate packaging of the genome-derived single exon microarray and the bioinformatic information is not required.

The amount of amplifiable probe material should be sufficient to permit at least one amplification 10 sufficient for subsequent hybridization assay.

Although the use of high density genome-derived microarrays on solid planar substrates is presently a preferred approach for the physical confirmation and characterization of the expression of sequences predicted 15 to encode protein, other types of microarrays (as herein defined) can also be used.

Furthermore, as earlier mentioned, experimental verification of the function predicted from genomic sequence in process 200 can be bioinformatic, rather than, 20 or additional to, physical verification.

For example, where the function desired to be identified is protein coding, the predicted ORFs can be compared bioinformatically to sequences known or suspected of being expressed.

Thus, the sequences output from process 300 (or process 200), can be used to query expression databases, such as EST databases, SNP ("single nucleotide polymorphism") databases, known cDNA and mRNA sequences, SAGE ("serial analysis of gene expression") databases, and 30 more generalized sequence databases that allow query for expressed sequences. Such query can be done by any sequence query algorithm, such as BLAST ("basic local alignment search tool"). The results of such query including information on identical sequences and . 35 information on nonidentical sequences that have diffuse or

focal regions of sequence homology to the query sequence — can then be passed directly to process 500, or used to inform analyses subsequently undertaken in process 200, process 300, or process 400.

or bioinformatic assay in process 400, is passed to process 500 where it is usefully related to the sequence data itself, a process colloquially termed "annotation". Such annotation can be done using any technique that usefully relates the functional information to the sequence, as, for example, by incorporating the functional data into the record itself, by linking records in a hierarchical or relational database, by linking to external databases, or by a combination thereof. Such database techniques are well within the skill in the art.

The annotated sequence data can be stored locally, uploaded to genomic sequence database 100, and/or displayed 800.

The methods and apparatus of the present

20 invention rapidly produce functional information from
genomic sequence. Coupled with the escalating pace at
which sequence now accumulates, the rapid pace of sequence
annotation produces a need for methods of displaying the
information in meaningful ways.

FIG. 3 shows visual display 80 presenting a single genomic sequence annotated according to the present invention. Because of its nominal resemblance to artistic works of Piet Mondrian, visual display 80 is alternatively described herein as a "Mondrian".

aligned with respect to the genomic sequence being annotated (hereinafter, the "annotated sequence"). Given the number of nucleotides typically represented in an annotated sequence, representation of individual

35 nucleotides would rarely be readable in hard copy output of

display 80. Typically, therefore, the annotated sequence is schematized as rectangle 89, extending from the left border of display 80 to its right border. By convention herein, the left border of rectangle 89 represents the first nucleotide of the sequence and the right border of rectangle 89 represents the last nucleotide of the sequence.

As further discussed below, however, the Mondrian visual display of annotated sequence can serve as a convenient graphical user interface for computerized representation, analysis, and query of information stored electronically. For such use, the individual nucleotides can conveniently be linked to the X axis coordinate of rectangle 89. This permits the annotated sequence at any point within rectangle 89 readily to be viewed, either automatically — for example, by time-delayed appearance of a small overlaid window upon movement of a cursor or other pointer over rectangle 89 — or through user intervention, as by clicking a mouse or other pointing device at a point in rectangle 89.

Visual display 80 is generated after user specification of the genomic sequence to be displayed. Such specification can consist of or include an accession number for a single clone (e.g., a single BAC accessioned into GenBank), wherein the starting and stopping nucleotides are thus absolutely identified, or alternatively can consist of or include an anchor or fulcrum point about which a chosen range of sequence is anchored, thus providing relative endpoints for the sequence to be displayed. For example, the user can anchor such a range about a given chromosomal map location, gene name, or even a sequence returned by query for similarity or identity to an input query sequence. When visual display 80 is used as a graphical user interface to computerized data, additional control over the first and

last displayed nucleotide will typically be dynamically selectable, as by use of standard zooming and/or selection tools.

Field 81 of visual display 80 is used to present 5 the output from process 200, that is, to present the bioinformatic prediction of those sequences having the desired function within the genomic sequence. Functional sequences are typically indicated by at least one rectangle 83 (83a, 83b, 83c), the left and right borders of which 10 respectively indicate, by their X-axis coordinates, the starting and ending nucleotides of the region predicted to have function.

Where a single bioinformatic method or approach identifies a plurality of regions having the desired 15 function, a plurality of rectangles 83 is disposed horizontally in field 81. Where multiple methods and/or approaches are used to identify function, each such method and/or approach can be represented by its own series of horizontally disposed rectangles 83, each such horizontally 20 disposed series of rectangles offset vertically from those representing the results of the other methods and approaches.

Thus, rectangles 83a in FIG. 3 represent the functional predictions of a first method of a first 25 approach for predicting function, rectangles 83b represent the functional predictions of a second method and/or second approach for predicting that function, and rectangles 83c represent the predictions of a third method and/or approach.

Where the function desired to be identified is protein coding, field 81 is used to present the bioinformatic prediction of sequences encoding protein. For example, rectangles 83a can represent the results from GRAIL or GRAIL II, rectangles 83b can represent the results 35 from GENEFINDER, and rectangles 83c can represent the

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results from DICTION.

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Optionally, and preferably, rectangles 83 collectively representing predictions of a single method and/or approach are identically colored and/or textured, and are distinguishable from the color and/or texture used for a different method and/or approach.

Alternatively, or in addition, the color, hue, density, or texture of rectangles 83 can be used further to report a measure of the bioinformatic reliability of the 10 prediction. For example, many gene prediction programs will report a measure of the reliability of prediction. Thus, increasing degrees of such reliability can be indicated, e.g., by increasing density of shading. Where display 80 is used as a graphical user interface, such 15 measures of reliability, and indeed all other results output by the program, can additionally or alternatively be made accessible through linkage from individual rectangles 83, as by time-delayed window ("tool tip" window), or by pointer (e.g., mouse) -activated link.

As earlier described, increased predictive reliability can be achieved by requiring consensus among methods and/or approaches to determining function. Thus, field 81 can include a horizontal series of rectangles 83 that indicate one or more degrees of consensus in 25 predictions of function.

Although FIG. 3 shows three series of horizontally disposed rectangles in field 81, display 80 can include as few as one such series of rectangles and as many as can discriminably be displayed, depending upon the 30 number of methods and/or approaches used to predict a given function.

Furthermore, field 81 can be used to show predictions of a plurality of different functions. However, the increased visual complexity occasioned by such 35 display makes more useful the ability of the user to select

a single function for display. When display 80 is used as a graphical user interface for computer query and analysis, such function can usefully be indicated and userselectable, as by a series of graphical buttons or tabs 5 (not shown in FIG. 3).

Rectangle 89 is shown in FIG. 3 as including interposed rectangle 84. Rectangle 84 represents the portion of annotated sequence for which predicted functional information has been assayed physically, with 10 the starting and ending nucleotides of the assayed material indicated by the X axis coordinates of the left and right borders of rectangle 84. Rectangle 85, with optional inclusive circles 86 (86a, 86b, and 86c) displays the results of such physical assay.

Although a single rectangle 84 is shown in FIG. 3, physical assay is not limited to just one region of annotated genomic sequence. It is expected that an increasing percentage of regions predicted to have function by process 200 will be assayed physically, and that display 20 80 will accordingly, for any given genomic sequence, have an increasing number of rectangles 84 and 85, representing an increased density of sequence annotation.

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Where the function desired to be identified is protein coding, rectangle 84 identifies the sequence of the 25 probe used to measure expression. In embodiments of the present invention where expression is measured using genome-derived single exon microarrays, rectangle 84 identifies the sequence included within the probe immobilized on the support surface of the microarray. 30 noted supra, such probe will often include a small amount of additional, synthetic, material incorporated during amplification and designed to permit reamplification of the probe, which sequence is typically not shown in display 80.

Rectangle 87 is used to present the results of 35 bioinformatic assay of the genomic sequence. For example,

where the function desired to be identified is protein coding, process 400 can include bioinformatic query of expression databases with the sequences predicted in process 200 to encode exons. And as earlier discussed,

5 because bioinformatic assay presents fewer constraints than does physical assay, often the entire output of process 200 can be used for such assay, without further subsetting thereof by process 300. Therefore, rectangle 87 typically need not have separate indicators therein of regions

10 submitted for bioinformatic assay; that is, rectangle 87 typically need not have regions therein analogous to rectangles 84 within rectangle 89.

Rectangle 87 as shown in FIG. 3 includes smaller rectangles 880 and 88. Rectangles 880 indicate regions
15 that returned a positive result in the bioinformatic assay, with rectangles 88 representing regions that did not return such positive results. Where the function desired to be predicted and displayed is protein coding, rectangles 880 indicate regions of the predicted exons that identify
20 sequence with significant similarity in expression databases, such as EST, SNP, SAGE databases, with rectangles 88 indicating genes novel over those identified in existing expression data bases.

Rectangles 880 can further indicate, through color, shading, texture, or the like, additional information obtained from bioinformatic assay.

For example, where the function assayed and displayed is protein coding, the degree of shading of rectangles 880 can be used to represent the degree of sequence similarity found upon query of expression databases. The number of levels of discrimination can be as few as two (identity, and similarity, where similarity has a user-selectable lower threshold). Alternatively, as many different levels of discrimination can be indicated as can visually be discriminated.

Where display 80 is used as a graphical user interface, rectangles 880 can additionally provide links directly to the sequences identified by the query of expression databases, and/or statistical summaries thereof.

5 As with each of the precedingly-discussed uses of display 80 as a graphical user interface, it should be understood that the information accessed via display 80 need not be resident on the computer presenting such display, which often will be serving as a client, with the linked information resident on one or more remotely located servers.

Rectangle 85 displays the results of physical assay of the sequence delimited by its left and right borders.

15 Rectangle 85 can consist of a single rectangle, thus indicating a single assay, or alternatively, and increasingly typically, will consist of a series of rectangles (85a, 85b, 85c) indicating separate physical assays of the same sequence.

20 Where the function assayed is gene expression, and where gene expression is assayed as herein described using simultaneous two-color fluorescent detection of hybridization to genome-derived single exon microarrays, individual rectangles 85 can be colored to indicate the 25 degree of expression relative to control. Conveniently, shades of green can be used to depict expression in the sample over control values, and shades of red used to depict expression less than control, corresponding to the spectra of the Cy3 and Cy5 dyes conventionally used for 30 respective labeling thereof. Additional functional information can be provided in the form of circles 86 (86a, 86b, 86c), where the diameter of the circle can be used to indicate expression intensity. As discussed infra, such relative expression (expression ratios) and absolute 35 expression (signal intensity) can be expressed using

normalized values.

Where display 80 is used as a graphical user interface, rectangle 85 can be used as a link to further information about the assay. For example, where the assay is one for gene expression, each rectangle 85 can be used to link to information about the source of the hybridized mRNA, the identity of the control, raw or processed data from the microarray scan, or the like.

gene prediction and gene expression for a hypothetical BAC, showing conventions used in the Examples presented infra.

BAC sequence ("Chip seq.") 89 is presented, with the physically assayed region thereof (corresponding to rectangle 84 in FIG. 3) shown in white. Algorithmic gene predictions are shown in field 81, with predictions by GRAIL shown, predictions by GENEFINDER, and predictions by DICTION shown. Within rectangle 87, regions of sequence that, when used to query expression databases, return identical or similar sequences ("EST hit") are shown as white rectangles (corresponding to rectangles 880 in FIG. 3), gray indicates low homology, and black indicates unknowns (where black and gray would correspond to rectangles 88 in FIG. 3).

Although FIGS. 3 and 4 show a single stretch of sequence, uninterrupted from left to right, longer sequences are usefully represented by vertical stacking of such individual Mondrians, as shown in FIGS. 9 and 10.

Single Exon Probes Useful For Measuring Gene Expression

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The methods and apparatus of the present invention rapidly produce functional information from genomic sequence. Where the function to be identified is protein coding, the methods and apparatus of the present invention rapidly identify and confirm the expression of

portions of genomic sequence that function to encode protein. As a direct result, the methods and apparatus of the present invention rapidly yield large numbers of single-exon nucleic acid probes, the majority from previously unknown genes, each of which is useful for measuring and/or surveying expression of a specific gene in one or more tissues or cell types.

It is, therefore, another aspect of the present invention to provide genome-derived single exon nucleic acid probes useful for gene expression analysis, and particularly for gene expression analysis by microarray.

Using the methods and genome-derived single-exon microarrays of the present invention, we have for example readily identified a large number of unique ORFs from human genomic sequence. Using single exon probes that encompass these ORFs, we have demonstrated, through microarray hybridization analysis, the expression of 9,980 of these ORFs in heart.

As would immediately be appreciated by one of skill in the art, each single exon probe having demonstrable expression in heart is currently available for use in measuring the level of its ORF's expression in heart.

Diseases of the heart and vascular system are a

25 significant cause of human morbidity and mortality.

Increasingly, genetic factors are being found that

contribute to predisposition, onset, and/or aggressiveness

of most, if not all, of these diseases. Although mutations

in single genes have on occasion been identified as

30 causative, these disorders are for the most part believed

to have polygenic etiologies.

For example, cardiovascular disease (CVD), which includes coronary heart disease, stroke, and peripheral arterial vascular disease, is the leading cause of death in the United States and other developed countries. In

developing regions, coronary heart disease and stroke are ranked second and third, respectively, as causes of mortality. In the United States alone, about 1 million deaths (about 42% of total deaths per year) result from CVD each year. CVD is also a significant cause of morbidity, with about 1.5 million people suffering myocardial infarction, and about 500,000 suffering strokes in the United States each year. With risk for CVD increasing with age, and an increasingly aging population, CVD will continue to be a major health problem into the future.

CVD is caused by arterial lesions that begin as fatty streaks, which consist of lipid-laden foam cells, and develop into fibrous plaques. The atherosclerotic plaque may grow slowly, and over several decades may produce a severe stenosis or result in arterial occlusion. Some plaques are stable, but other, more unstable, ones may rupture and induce thrombosis. The thrombi may embolize, rapidly occluding the lumen and leading to myocardial infarction or acute ischemic syndrome.

Risk factors for CVD include age and gender. 20 addition, a family history of CVD significantly increases risk, indicating a genetic basis for development of this disease complex. Obesity, especially truncal obesity, the cause of which is suspected to be genetic, is yet another 25 risk factor for CVD. Familial disorders such as hyperlipidemia, hypoalphalipoproteinemia, hypertriglyceridemia, hypercholesterolemia, hyperinsulinemia, homocystinuria, and dysbetalipoproteinemia, all of which lead to lipid or 30 lipoprotein abnormalities, can predispose one to the development of CVD. Both insulin-dependent and noninsulin-dependent diabetes mellitus, both of which have genetic components, have been also linked to the development of atherosclerosis.

35 The literature is replete with evidence for

genetic causes of cardiovascular diseases. For example, studies by Allayee et al., Am. J. Hum. Genet. 63:577-585(1998), indicated a genetic association between familial combined hyperlipidemia (FCHL) and small dense LDL particles. The studies also concluded that the genetic determinants for LDL particle size are shared, at least in part, among FCHL families and the more general population at risk for CVD. Juo et al., Am. J. Hum. Genet. 63: 586-594 (1998) demonstrated that small, dense LDL particles and elevated apolipoprotein B levels, both of which are commonly found in members of FCHL families, share a common major gene plus individual polygenic components.

The common major gene was estimated to explain 37% of the variants of adjusted LDL particle size and 23% of the variants of adjusted apoB levels.

The atherogenic lipoprotein phenotype (ALP) is a common heritable trait, symptoms of which include a prevalence of small, dense LDL particles, increased levels of triglyceride-rich lipoproteins, reduced levels of high density lipoprotein, and increased risk of CVD, particularly myocardial infarction. Both Nishina et al., Proc. Nat. Acad. Sci. 89: 708-712 (1992) and Rotter et al., Am. J. Hum. Genet. 58: 585-594(1996) demonstrated linkage between ALP and the LDLR locus. Rotter et al., supra, also reported linkage to the CETP locus on chromosome 16 and to the SOD1 locus on chromosome 6, and possibly also to the APOA1/APOC3/APOA4 cluster on chromosome 11.

Mutations in genes identified as components of lipid metabolism, e.g., apolipoprotein E (apoE) and LDL receptor (LDLR), have been shown to be associated with predisposition to the development of CVD. For example, several apoE variants had been found to be associated with familial dysbetalipoproteinemia, characterized by elevated plasma cholesterol and triglyceride levels and an increased risk for atherosclerosis (de Knijff et al., Mutat 4: 178-

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194 (1994)). Mutations in the LDLR gene have been associated with the familial hypercholesterolemia, an autosomal dominant disorder characterized by elevation of serum cholesterol bound to low density lipoprotein (LDL), that can lead to increased susceptibility to CVD.

To date, mutations in numerous genes have been shown to be associated with increased CVD susceptibility. However, the identified genetic associations are believed not to account for all genetic contributions to CVD.

As yet another example, hypertension is a major health problem because of its high prevalence and its association with increased risk of CVD. Approximately 25% of all adults and over 60% of persons older than 60 years in the United States have high blood pressure.

15 Arterial or systemic hypertension is diagnosed when the average of two or more diastolic BP measurements on at least two subsequent visits is 90 mm Hg or more, or when the average of multiple systolic BP readings on two or more subsequent visits is consistently greater than 140 mm
20 Hg. Pulmonary hypertension is defined as pressure within the pulmonary arterial system elevated above the normal range; pulmonary hypertension may lead to right ventricle (RV) failure.

Hypertension, together with other cardiovascular risk factors, leads to atherosclerosis and other forms of CVD, primarily by damaging the vascular endothelium. In more than 40% of the U.S. population, hypertension is accompanied by hyperlipidemia and leads to the development of atherosclerotic plaques. In the absence of hyperlipidemia, intimal thickening occurs. Non-atherosclerotic hypertension-induced vascular damage can

lead to stroke or heart failure.

Familial diseases associated with secondary hypertension include familial renal disease, polycystic kidney disease, medullary thyroid cancer, pheochromocytoma,

and hyperparathyroidism. Hypertension is also twice as common in patients with diabetes mellitus.

More than 95% of all hypertension cases are essential hypertension, that is, lack identifiable

5 antecedent clinical cause. Essential hypertension shows clustering in families and can result from a variety of genetic diseases. In most cases, high blood pressure results from a complex interaction of factors with both genetic and environmental components. The recent search for genes that contribute to the development of essential hypertension has shown that the disorder is polygenic in origin. However, with several exceptions (such as angiotensinogen, angiotensin receptor-1, beta-3 subunit of guanine nucleotide-binding protein, tumor necrosis factor receptor-2, and "-adducin), the particular genes involved are still being sought.

Susceptibility loci for essential hypertension have been mapped to chromosomes 17 and 15q. Hasstedt et al., Am. J. Hum. Genet. 43: 14-22 (1988) measured red cell sodium in 1,800 normotensive members of 16 Utah pedigrees ascertained through hypertensive or normotensive probands, siblings with early stroke death, or brothers with early coronary disease, and suggested that red blood cell sodium was determined by 4 alleles at a single locus. This major locus was thought to explain 29% of the variance in red cell sodium, and polygenic inheritance explained another 54.6%. A higher frequency of the high red blood cell sodium genotype in pedigrees in which the proband was hypertensive rather than normotensive provided evidence that this major locus increases susceptibility to hypertension.

From a study of systolic blood pressure in 278 pedigrees, Perusse et al., Am. J. Hum. Genet. 49: 94-105 (1991) reported that variability in systolic blood pressure is likely influenced by allelic variation of a single gene, with gender and age dependence. They also suggested that a

single gene may be associated with a steeper increase of blood pressure with age among males and females.

There is strong evidence, however, for additional as yet uncharacterized, hypertension-associated loci on other chromosomes.

For example, Xu et al., Am. J. Hum. Genet. 64: 1694-1701 (1999) carried out a systematic search for chromosomal regions containing genes that regulate blood pressure by scanning the entire autosomal genome using 367 polymorphic markers. Because of the sampling design, the number of sib pairs, and the availability of genotyped parents, this study represented one of the most powerful of its kind. Although no regions achieved a 5% genomewide significance level, maximum lod scores were greater than 2.0 for regions of chromosomes 3, 11, 15, 16, and 17.

As another example, cardiac arrhythmias account for several thousand deaths each year. Arrhythmias such as ventricular fibrillation, which causes more than 300,000 sudden deaths annually in the United States alone,

20 encompass a multitude of disorders. Another type of arrhythmia, idiopathic dilated cardiomyopathy, of which familial dilated cardiomyopathy accounts for 20-25%, is responsible for more than 10,000 deaths in the United States annually and is the predominant indication for cardiac transplantation.

Cardiac arrhythmias can be divided into bradyarrhythmias (slowed rhythms) or tachyarrhythmias (speeded rhythms). Bradyarrhythmias result from abnormalities of intrinsic automatic behavior or conduction, primarily within the atrioventricular node and the His-Purkinje's network. Tachyarrhythmias are caused by altered automaticity, reentry, or triggered automaticity.

Bradyarrhythmias arising from suspected polygenic disorders include Long QT syndrome 4, atrioventricular block, familial sinus node disease, progressive cardiac

conduction defect, and familial cardiomyopathy.

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Tachyarrhythmias with possible underlying polygenic causes include familial ventricular tachycardia, Wolff-Parkinson-White syndrome, familial arrhythmogenic right ventricular 5 dysplasia, heart-hand syndrome V, Mal de Meleda, familial ventricular fibrillation, and familial noncompaction of left ventricular myocardium.

For some of the arrhythmias, one or more of the causative genes have been identified.

For example, atrioventricular block has been associated with mutations in the SCN5A gene, as well as mutations in a locus mapped to 19q13. Studies have shown linkage of familial sinus node disease to a marker on 10g22-g24. Familial ventricular tachycardia has been 15 linked to mutations in genes encoding the G protein subunit

alpha-i2 (GNAI1), and/or related genes. Examination of families with Wolff-Parkinson-White syndrome suggest an autosomal dominant pattern of inheritance and evidence of linkage of the disorder to DNA markers on band 7q3.

20 Linkage analysis shows strong evidence for localization of a gene for Mal de Meleda disease on 8qter. Familial ventricular fibrillation can be caused by mutations in the cardiac sodium channel gene SCN5A. Familial noncompaction of left ventricular myocardium has been linked to mutations 25 in the gene encoding tafazzin (TAZ), or in the FK506-

binding protein 1A gene (FKBP1A).

Familial dilated cardiomyopathy is characterized by an autosomal dominant pattern of inheritance with agerelated penetrance. The linkage of familial dilated 30 cardiomyopathy to several loci indicate that it is polygenic. These loci include CMD1A on 1p11-q11, CMD1B on 9q13, CMD1C on 10q21, CMD1D on 1q32, CMD1E on 3p, CMD1F on 6q, CMD1G on 2q31, CMD1H on 2q14-q22, and CMD1I, which results from mutation in the DES gene on 2q35.

In addition, cardiomyopathy can also be caused by 35

mutations in the ACTC gene, the cardiac beta-myosin heavy chain gene (MYH7), or the cardiac troponin T gene.

Familial arrhythmogenic right ventricular dysplasia is inherited as an autosomal dominant with reduced penetrance and is one of the major genetic causes of juvenile sudden death. It is estimated that the prevalence of familial arrhythmogenic right ventricular dysplasia ranges from 6 per 10,000 in the general population to 4.4 per 1,000 in some areas.

Several loci for familial arrhythmogenic right ventricular dysplasia have been mapped indicating that this disease is also polygenic in nature. These loci include ARVD1 on 14q23-q24, ARVD2 on 1q42-q43, ARVD3 on 14q12-q22, ARVD4 on 2q32.1-q32.3, ARVD5 on 3p23, and ARVD6 on 10p14-p12.

Progressive cardiac conduction defect (PCCD), also called Lenegre-Lev disease, is one of the most common cardiac conduction diseases. It is characterized by progressive alteration of cardiac conduction through the His-Purkinje system with right or left bundle branch block and widening of QRS complexes, leading to complete atrioventricular block and ultimately causing syncope and sudden death. It represents the major cause of pacemaker implantation in the world (0.15 implantations per 1,000 inhabitants per year in developed countries). The cause of PCCD is unknown but familial cases with right bundle branch block have been reported suggesting that at least some cases are of genetic origin. Reports have linked PCCD to HB1 on 19q13.3, and to mutations in the SCN5A gene (Schott et al., Nature Genet. 23: 20-21 (1999)).

As yet a further example, congenital heart disease occurs at a rate of 8 per 1000 live births, which corresponds to approximately 32,000 infants with newly diagnosed congenital heart disease each year in the United States. Twenty percent of infants with congenital heart

disease die within the first year of life. Approximately 80% of the first-year survivors live to reach adulthood. Congenital heart disease also has economic impact due to the estimated 20,000 surgical procedures performed to correct circulatory defects in these patients. The estimated number of adults with congenital heart disease in the United States is currently about 900,000.

In 90% of patients, congenital heart disease is attributable to multifactorial inheritance. Only 5-10% of malformations are due to primary genetic factors, which are either chromosomal or a result of a single mutant gene.

The most common congenital heart disease found in adults is bicuspid aortic valve. This defect occurs in 2% of the general population and accounts for approximately 50% of operated cases of aortic stenosis in adults. Atrial septal defect is responsible for 30-40% of congenital heart disease seen in adults. The most common congenital cardiac defect observed in the pediatric population is ventricular septal defect, which accounts for 15-20% of all congenital lesions. Tetralogy of Fallot is the most common cyanotic congenital anomaly observed in adults. Other congenital heart diseases include Eisenmenger's syndrome, patent ductus arteriosus, pulmonary stenosis, coarctation of the aorta, transposition of the great arteries, tricuspid atresia, univentricular heart, Ebstein's anomaly, and double-outlet right ventricle.

A number of studies have identified putative genetic loci associated with one or more congenital heart diseases.

Congenital heart disease affects more than 40% of all Down syndrome patients. The candidate chromosomal region containing the putative gene or genes for congenital heart disease associated with Down syndrome is 21q22.2-q22.3, between ETS2 and MX1.

DiGeorge syndrome (DGS) is characterized by

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several symptoms including outflow tract defects of the heart such as teratology of Fallot. Most cases result from a deletion of chromosome 22q11.2 (the DiGeorge syndrome chromosome region, or DGCR). The 22q11 deletion is the second most common cause of congenital heart disease after Down syndrome. Several genes are lost in this deletion including the putative transcription factor TUPLE1. This deletion is associated with a variety of phenotypes, e.g., Shprintzen syndrome; conotruncal anomaly face (or Takao syndrome); and isolated outflow tract defects of the heart including Tetralogy of Fallot, truncus arteriosus, and interrupted aortic arch.

Whereas 90% of cases of DGS may now be attributed to a 22q11 deletion, other associated chromosome defects

15 have been identified. For example, Greenberg et al., Am.

J. Hum. Genet. 43:605-611 (1988), reported 1 case of DGS

with del10p13 and one with a 18q21.33 deletion. Fukushima et al., Am. J. Hum. Genet. 51 (suppl.):A80 (1992) reported linkage with a deletion of 4q21.3-q25. Gottlieb et al.,

20 Am. J. Hum. Genet. 62: 495-498 (1998) concluded that the deletion of more than 1 region on 10p could be associated with the DGS phenotype. The association of the DiGeorge syndrome with at least 2 and possibly more chromosomal locations suggests strongly the involvement of several

25 genes in this disease.

Digilio et al., J. Med. Genet. 34: 188-190 (1997), calculated empiric risk figures for recurrence of isolated Tetralogy of Fallot in families after exclusion of del(22q11), and concluded that gene(s) different from those located on 22q11 must be involved in causing familial aggregation of nonsyndromic Tetralogy of Fallot. Johnson et al., Am. J. Med. Genet. (1997) conducted a cytogenetic evaluation of 159 cases of Tetralogy of Fallot. They reported that a del(22q11) was identified in 14% who underwent fluorescence in situ hybridization (FISH) testing

with the N25 cosmid probe.

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Other congenital heart disease are also suspected to be of polygenic origin. For example, Holmes et al., Birth Defects Orig. Art. Ser. X(4): 228-230 (1974)

5 described familial clustering of hypoplastic left heart syndrome in siblings consistent with multifactorial causation.

Other significant diseases of the heart and vascular system are also believed to have a genetic, 10 typically polygenic, etiological component. These diseases include, for example, hypoplastic left heart syndrome, cardiac valvular dysplasia, Pfeiffer cardiocranial syndrome, oculofaciocardiodental syndrome, Kapur-Toriello syndrome, Sonoda syndrome, Ohdo Blepharophimosis syndrome, 15 heart-hand syndrome, Pierre-Robin syndrome, Hirschsprung disease, Kousseff syndrome, Grange occlusive arterial syndrome, Kearns-Sayre syndrome, Kartagener syndrome, Alagille syndrome, Ritscher-Schinzel syndrome, Ivemark syndrome, Young-Simpson syndrome, hemochromatosis, 20 Holzgreve syndrome, Barth syndrome, Smith-Lemli-Opitz syndrome, glycogen storage disease, Gaucher-like disease, Fabry disease, Lowry-Maclean syndrome, Rett syndrome, Opitz syndrome, Marfan syndrome, Miller-Dieker lissencephaly syndrome, mucopolysaccharidosis, Bruada syndrome, 25 humerospinal dysostosis, Phaver syndrome, McDonough syndrome, Marfanoid hypermobility syndrome, atransferrinemia, Cornelia de Lange syndrome, Leopard syndrome, Diamond-Blackfan anemia, Steinfeld syndrome, progeria, and Williams-Beuren syndrome.

The human genome-derived single exon nucleic acid probes and microarrays of the present invention are useful for predicting, diagnosing, grading, staging, monitoring and prognosing diseases of human heart and vascular system, particularly those diseases with polygenic etiology. With each of the single exon probes described herein shown to be

expressed at detectable levels in human heart, and with about 2/3 of the probes identifying novel genes, the single exon microarrays of the present invention provide exceptionally high informational content for such studies.

For example, diagnosis (including differential diagnosis among clinically indistinguishable disorders), staging, and/or grading of a disease can be based upon the quantitative relatedness of a patient gene expression profile to one or more reference expression profiles known to be characteristic of a given heart or vascular disease, or to specific grades or stages thereof.

In one embodiment, the patient gene expression profile is generated by hybridizing nucleic acids obtained directly or indirectly from transcripts expressed in the patient's heart or vascular tissues to the genome-derived single exon microarray of the present invention. Reference profiles are obtained similarly by hybridizing nucleic acids obtained directly or indirectly from transcripts expressed in heart or vascular tissue of individuals with known disease. Methods for quantitatively relating gene expression profiles, without regard to the function of the protein encoded by the gene, are disclosed in WO 99/58720, incorporated herein by reference in its entirety.

In another approach, the genome-derived single

exon probes and microarrays of the present invention can be
used to interrogate genomic DNA, rather than pools of
expressed message; this latter approach permits
predisposition to and/or prognosis of heart or vascular
disease to be assessed through the massively parallel

determination of altered copy number, deletion, or mutation
in the patient's genome of exons known to be expressed in
human heart. The algorithms set forth in WO 99/58720 can
be applied to such genomic profiles without regard to the
function of the protein encoded by the interrogated gene.

The utility is specific to the probe; at

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sufficiently high hybridization stringency, which stringencies are well known in the art - see Ausubel et al. and Maniatis et al. - each probe reports the level of expression of message specifically containing that ORF.

It should be appreciated, however, that the probes of the present invention, for which expression in the heart has been demonstrated are useful for both measurement in the heart and for survey of expression in other tissues.

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Significant among such advantages is the presence of probes for novel genes.

As mentioned above and further detailed in Examples 1 and 2, the methods described enable ORFs which are not present in existing expression databases to be identified. And the fewer the number of tissues in which the ORF can be shown to be expressed, the more likely the ORF will prove to be part of a novel gene: as further discussed in Example 2, ORFs whose expression was measurable in only a single of the tested tissues were 20 represented in existing expression databases at a rate of only 11%, whereas 36% of ORFs whose expression was measurable in 9 tissues were present in existing expression databases, and fully 45% of those ORFs expressed in all ten tested tissues were present in existing expressed sequence databases.

Either as tools for measuring gene expression or tools for surveying gene expression, the genome-derived single exon probes of the present invention have significant advantages over the cDNA or EST-based probes that are currently available for achieving these utilities.

The genome-derived single exon probes of the present invention are useful in constructing genome-derived single exon microarrays; the genome-derived single exon microarrays, in turn, are useful devices for measuring and for surveying gene expression in the human.

Gene expression analysis using microarrays — conventionally using microarrays having probes derived from expressed message — is well-established as useful in the biological research arts (see Lockhart et al. Nature 405, 827-836).

Microarrays have been used to determine gene expression profiles in cells in response to drug treatment (see, for example, Kaminski et al., "Global Analysis of Gene Expression in Pulmonary Fibrosis Reveals Distinct 10 Programs Regulating Lung Inflammation and Fibrosis," Proc. Natl. Acad. Sci. USA 97(4):1778-83 (2000); Bartosiewicz et al., "Development of a Toxicological Gene Array and Quantitative Assessment of This Technology, " Arch. Biochem. Biophys. 376(1):66-73 (2000)), viral infection (see for 15 example, Geiss et al., "Large-scale Monitoring of Host Cell Gene Expression During HIV-1 Infection Using cDNA Microarrays," Virology 266(1):8-16 (2000)) and during cell processes such as differentiation, senescence and apoptosis (see, for example, Shelton et al., "Microarray Analysis of 20 Replicative Senescence, " Curr. Biol. 9(17):939-45 (1999); Voehringer et al., "Gene Microarray Identification of Redox and Mitochondrial Elements That Control Resistance or Sensitivity to Apoptosis," Proc. Natl. Acad. Sci. USA 97(6):2680-5 (2000)).

Microarrays have also been used to determine abnormal gene expression in diseased tissues (see, for example, Alon et al., "Broad Patterns of Gene Expression Revealed by Clustering Analysis of Tumor and Normal Colon Tissues Probed by Oligonucleotide Arrays," Proc. Natl.

30 Acad. Sci. USA 96(12):6745-50 (1999); Perou et al., "Distinctive Gene Expression Patterns in Human Mammary Epithelial Cells and Breast Cancers, Proc. Natl. Acad. Sci. USA 96(16):9212-7 (1999); Wang et al., "Identification of Genes Differentially Over-expressed in Lung Squamous Cell Carcinoma Using Combination of cDNA Subtraction and

Microarray Analysis, "Oncogene 19(12):1519-28 (2000);
Whitney et al., "Analysis of Gene Expression in Multiple
Sclerosis Lesions Using cDNA Microarrays," Ann. Neurol.
46(3):425-8 (1999)), in drug discovery screens (see, for
example, Scherf et al., "A Gene Expression Database for the
Molecular Pharmacology of Cancer," Nat. Genet. 24(3):236-44
(2000)) and in diagnosis to determine appropriate treatment
strategies (see, for example, Sgroi et al., "In vivo Gene
Expression Profile Analysis of Human Breast Cancer
Progression," Cancer Res. 59(22):5656-61 (1999)).

In microarray-based gene expression screens of pharmacological drug candidates upon cells, each probe provides specific useful data. In particular, it should be appreciated that even those probes that show no change in expression are as informative as those that do change, serving, in essence, as negative controls.

For example, where gene expression analysis is used to assess toxicity of chemical agents on cells, the failure of the agent to change a gene's expression level is evidence that the drug likely does not affect the pathway of which the gene's expressed protein is a part.

Analogously, where gene expression analysis is used to assess side effects of pharmacological agents — whether in lead compound discovery or in subsequent screening of lead compound derivatives — the inability of the agent to alter a gene's expression level is evidence that the drug does not affect the pathway of which the gene's expressed protein is a part.

WO 99/58720 provides methods for quantifying the relatedness of a first and second gene expression profile and for ordering the relatedness of a plurality of gene expression profiles. The methods so described permit useful information to be extracted from a greater percentage of the individual gene expression measurements from a microarray than methods previously used in the art.

Other uses of microarrays are described in Gerhold et al., Trends Biochem. Sci. 24(5):168-173 (1999) and Zweiger, Trends Biotechnol. 17(11):429-436 (1999); Schena et al.

5 The invention particularly provides genomederived single-exon probes known to be expressed in heart.

The individual single exon probes can be provided in the form of substantially isolated and purified nucleic acid, typically, but not necessarily, in a quantity sufficient to perform a hybridization reaction.

Such nucleic acid can be in any form directly hybridizable to the message that contains the probe's ORF, such as double stranded DNA, single-stranded DNA complementary to the message, single-stranded RNA complementary to the message, or chimeric DNA/RNA molecules so hybridizable. The nucleic acid can alternatively or additionally include either nonnative nucleotides, alternative internucleotide linkages, or both, so long as complementary binding can be obtained. For example, probes can include phosphorothioates, methylphosphonates, morpholino analogs, and peptide nucleic acids (PNA), as are described, for example, in U.S. Patent Nos. 5,142,047; 5,235,033; 5,166,315; 5,217,866; 5,184,444; 5,861,250.

Usefully, however, such probes are provided in a form and quantity suitable for amplification, where the amplified product is thereafter to be used in the hybridization reactions that probe gene expression.

Typically, such probes are provided in a form and quantity suitable for amplification by PCR or by other well known amplification technique. One such technique additional to PCR is rolling circle amplification, as is described, inter alia, in U.S. Patent Nos. 5,854,033 and 5,714,320 and international patent publications WO 97/19193 and WO 00/15779. As is well understood, where the probes are to be provided in a form suitable for amplification, the

range of nucleic acid analogues and/or internucleotide linkages will be constrained by the requirements and nature of the amplification enzyme.

Where the probe is to be provided in form

5 suitable for amplification, the quantity need not be
sufficient for direct hybridization for gene expression
analysis, and need be sufficient only to function as an
amplification template, typically at least about 1, 10 or
100 pg or more.

Description Each discrete amplifiable probe can also be packaged with amplification primers, either in a single composition that comprises probe template and primers, or in a kit that comprises such primers separately packaged therefrom. As earlier mentioned, the ORF-specific 5' primers used for genomic amplification can have a first common sequence added thereto, and the ORF-specific 3' primers used for genomic amplification can have a second, different, common sequence added thereto, thus permitting, in this embodiment, the use of a single set of 5' and 3' primers to amplify any one of the probes. The probe composition and/or kit can also include buffers, enzyme, etc., required to effect amplification.

As mentioned earlier, when intended for use on a genome-derived single exon microarray of the present

25 invention, the genome-derived single exon probes of the present invention will typically average at least about 100, 200, 300, 400 or 500 bp in length, including (and typically, but not necessarily centered about) the ORF. Furthermore, when intended for use on a genome-derived

30 single exon microarray of the present invention, the genome-derived single exon probes of the present invention will typically not contain a detectable label.

When intended for use in solution phase hybridization, however — that is, for use in a hybridization reaction in which the probe is not first

bound to a support substrate (although the target may indeed be so bound) - length constraints that are imposed in microarray-based hybridization approaches will be relaxed, and such probes will typically be labeled.

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In such case, the only functional constraint that dictates the minimum size of such probe is that each such probe must be capable of specifically identifying in a hybridization reaction the exon from which it is drawn. theory, a probe of as little as 17 nucleotides is capable 10 of uniquely identifying its cognate sequence in the human genome. For hybridization to expressed message - a subset of target sequence that is much reduced in complexity as compared to genomic sequence - even fewer nucleotides are required for specificity.

Therefore, the probes of the present invention 15 can include as few as 20, 25 or 50 bp or ORF, or more. particular embodiments, the ORF sequences are given in SEQ ID NOS. 9,981 - 19,771, respectively, for probe SEQ ID NOS. 1 - 9,980. The minimum amount of ORF required to be 20 included in the probe of the present invention in order to provide specific signal in either solution phase or microarray-based hybridizations can readily be determined for each of ORF SEQ ID NOS. 9,981 - 19,771 individually by routine experimentation using standard high stringency 25 conditions.

Such high stringency conditions are described, inter alia, in Ausubel et al. and Maniatis et al. microarray-based hybridization, standard high stringency conditions can usefully be 50% formamide, 5X SSC, 0.2 µg/µl 30 poly(dA), 0.2 $\mu g/\mu l$ human c_ot1 DNA, and 0.5 % SDS, in a humid oven at 42°C overnight, followed by successive washes of the microarray in 1X SSC, 0.2% SDS at 55°C for 5 minutes, and then 0.1X SSC, 0.2% SDS, at 55°C for 20 minutes. For solution phase hybridization, standard high 35 stringency conditions can usefully be aqueous hybridization

at 65°C in 6X SSC. Lower stringency conditions, suitable for cross-hybridization to mRNA encoding structurally- and functionally-related proteins, can usefully be the same as the high stringency conditions but with reduction in 5 temperature for hybridization and washing to room temperature (approximately 25°C).

When intended for use in solution phase hybridization, the maximum size of the single exon probes of the present invention is dictated by the proximity of other expressed exons in genomic DNA: although each single exon probe can include intergenic and/or intronic material contiguous to the ORF in the human genome, each probe of the present invention will include portions of only one expressed exon.

Thus, each single exon probe will include no more than about 25 kb of contiguous genomic sequence, more typically no more than about 20 kb of contiguous genomic sequence, more usually no more than about 15 kb, even more usually no more than about 10 kb. Usually, probes that are 20 maximally about 5 kb will be used, more typically no more than about 3 kb.

15

It will be appreciated that the Sequence Listing appended hereto presents, by convention, only that strand of the probe and ORF sequence that can be directly 25 translated reading from 5' to 3' end. As would be well understood by one of skill in the art, single stranded probes must be complementary in sequence to the ORF as present in an mRNA; it is well within the skill in the art to determine such complementary sequence. It will further 30 be understood that double stranded probes can be used in both solution-phase hybridization and microarray-based hybridization if suitably denatured.

Thus, it is an aspect of the present invention to provide single-stranded nucleic acid probes that have 35 sequence complementary to those described herein above and

below, and double-stranded probes one strand of which has sequence complementary to the probes described herein.

The probes can, but need not, contain intergenic and/or intronic material that flanks the ORF, on one or both sides, in the same linear relationship to the ORF that the intergenic and/or intronic material bears to the ORF in genomic DNA. The probes do not, however, contain nucleic acid derived from more than one expressed ORF.

And when intended for use in solution

10 hybridization, the probes of the present invention can

11 usefully have detectable labels. Nucleic acid labels are

12 well known in the art, and include, inter alia, radioactive

13 labels, such as ³H, ³²P, ³³P, ³⁵S, ¹²⁵I, ¹³¹I; fluorescent

14 labels, such as Cy3, Cy5, Cy5.5, Cy7, SYBR®

15 Green and other labels described in Haugland,

Handbook of Fluorescent Probes and Research Chemicals, 7th
ed., Molecular Probes Inc., Eugene, OR (2000), or
fluorescence resonance energy transfer tandem conjugates
thereof; labels suitable for chemiluminescent and/or
20 enhanced chemiluminescent detection; labels suitable for
ESR and NMR detection; and labels that include one member
of a specific binding pair, such as biotin, digoxigenin, or
the like.

The probes, either in quantity sufficient for hybridization or sufficient for amplification, can be provided in individual vials or containers.

Alternatively, such probes can usefully be packaged as a plurality of such individual genome-derived single exon probes.

30 When provided as a collection of plural individual probes, the probes are typically made available in amplifiable form in a spatially-addressable ordered set, typically one per well of a microtiter dish. Although a 96 well microtiter plate can be used, greater efficiency is obtained using higher density arrays.

If, as earlier mentioned, the ORF-specific
5' primers used for genomic amplification had a first
common sequence added thereto, and the ORF-specific 3'
primers used for genomic amplification had a second,
different, common sequence added thereto, a single set of
5' and 3' primers can be used to amplify all of the probes
from the amplifiable ordered set.

Such collections of genome-derived single exon probes can usefully include a plurality of probes chosen for the common attribute of expression in the human heart.

In such defined subsets, typically at least 50, 60, 75, 80, 85, 90 or 95% or more of the probes will be chosen by their expression in the defined tissue or cell type.

The single exon probes of the present invention, as well as fragments of the single exon probes comprising selectively hybridizable portions of the probe ORF, can be used to obtain the full length cDNA that includes the ORF by (i) screening of cDNA libraries; (ii) rapid

20 amplification of cDNA ends ("RACE"); or (iii) other conventional means, as are described, inter alia, in Ausubel et al. and Maniatis et al.

It is another aspect of the present invention to provide genome-derived single exon nucleic acid microarrays useful for gene expression analysis, where the term "microarray" has the meaning given in the definitional section of this description, supra.

The invention particularly provides genome-derived single-exon nucleic acid microarrays comprising a plurality of probes known to be expressed in human heart. In preferred embodiments, the present invention provides human genome-derived single exon microarrays comprising a plurality of probes drawn from the group consisting of SEQ ID NOS.: 1 - 9,980.

When used for gene expression analysis, the

genome-derived single exon microarrays provide greater physical informational density than do the genome-derived single exon microarrays that have lower percentages of probes known to be expressed commonly in the tested tissue.

5 At a fixed probe density, for example, a given microarray surface area of the defined subset genome-derived single exon microarray can yield a greater number of expression measurements. Alternatively, at a given probe density, the same number of expression measurements can be obtained from a smaller substrate surface area. Alternatively, at a fixed probe density and fixed surface area, probes can be provided redundantly, providing greater reliability in signal measurement for any given probe. Furthermore, with a higher percentage of probes known to be expressed in the assayed tissue, the dynamic range of the detection means can be adjusted to reveal finer levels discrimination among the levels of expression.

Although particularly described with respect to their utility as probes of gene expression, particularly as probes to be included on a genome-derived single exon microarray, each of the nucleic acids having SEQ ID NOS.: 1 - 9,980 contains an open-reading frame, set forth respectively in SEQ ID NOS.: 9,981 - 19,771, that encodes a protein domain. Thus, each of SEQ ID NOS. 1 - 9,980 can be used, or that portion thereof in SEQ ID NOS. 9,981 - 19,771 used, to express a protein domain by standard in vitro recombinant techniques. See Ausubel et al. and Maniatis et al.

Additionally, kits are available commercially

that readily permit such nucleic acids to be expressed as protein in bacterial cells, insect cells, or mammalian cells, as desired (e.g., HAT Protein Expression & Purification System, ClonTech Laboratories, Palo Alto, CA; Adeno-XT Expression System, ClonTech Laboratories, Palo

Alto, CA; Protein Fusion & Purification (pMAL) System, New

England Biolabs, Beverley, MA)

Furthermore, shorter peptides can be chemically synthesized using commercial peptide synthesizing equipment and well known techniques. Procedures are described, inter alia, in Chan et al. (eds.), Fmoc Solid Phase Peptide Synthesis: A Practical Approach (Practical Approach Series, (Paper)), Oxford Univ. Press (March 2000) (ISBN: 0199637245); Jones, Amino Acid and Peptide Synthesis (Oxford Chemistry Primers, No 7), Oxford Univ. Press (August 1992) (ISBN: 0198556683); and Bodanszky, Principles of Peptide Synthesis (Springer Laboratory), Springer Verlag (December 1993) (ISBN: 0387564314).

It is, therefore, another aspect of the invention to provide peptides comprising an amino acid sequence

15 translated from SEQ ID NOS.: 9,981 - 19,771. Such amino acid sequences are set out in SEQ ID NOS: 19,772 - 29,119. Any such recombinantly-expressed or synthesized peptide of at least 8, and preferably at least about 15, amino acids, can be conjugated to a carrier protein and used to generate antibody that recognizes the peptide. Thus, it is a further aspect of the invention to provide peptides that have at least 8, preferably at least 15, consecutive amino acids.

The following examples are offered by way of illustration and not by way of limitation.

EXAMPLE 1

Preparation of Single Exon Microarrays from ORFs Predicted 30 in Human Genomic Sequence

Bioinformatics Results

All human BAC sequences in fewer than 10 pieces that had been accessioned in a five month period

35 immediately preceding this study were downloaded from

GenBank. This corresponds to ~2200 clones, totaling ~350 MB of sequence, or approximately 10% of the human genome.

After masking repetitive elements using the program CROSS_MATCH, the sequence was analyzed for open reading frames using three separate gene finding programs. The three programs predict genes using independent algorithmic methods developed on independent training sets: GRAIL uses a neural network, GENEFINDER uses a hidden Markoff model, and DICTION, a program proprietary to Genetics Institute, operates according to a different heuristic. The results of all three programs were used to create a prediction matrix across the segment of genomic DNA.

The three gene finding programs yielded a range of results. GRAIL identified the greatest percentage of genomic sequence as putative coding region, 2% of the data analyzed. GENEFINDER was second, calling 1%, and DICTION yielded the least putative coding region, with 0.8% of genomic sequence called as coding region.

The consensus data were as follows. GRAIL and GENEFINDER agreed on 0.7% of genomic sequence, GRAIL and DICTION agreed on 0.5% of genomic sequence, and the three programs together agreed on 0.25% of the data analyzed. That is, 0.25% of the genomic sequence was identified by all three of the programs as containing putative coding region.

ORFs predicted by any two of the three programs ("consensus ORFs") were assorted into "gene bins" using two criteria: (1) any 7 consecutive exons within a 25 kb window were placed together in a bin as likely contributing to a single gene, and (2) all ORFs within a 25 kb window were placed together in a bin as likely contributing to a single gene if fewer than 7 exons were found within the 25 kb window.

PCR ·

The largest ORF from each gene bin that did not span repetitive sequence was then chosen for amplification, as were all consensus ORFs longer than 500 bp. This method approximated one exon per gene; however, a number of genes were found to be represented by multiple elements.

Previously, we had determined that DNA fragments fewer than 250 bp in length do not bind well to the aminomodified glass surface of the slides used as support substrate for construction of microarrays; therefore, amplicons were designed in the present experiments to approximate 500 bp in length.

Accordingly, after selecting the largest ORF per gene bin, a 500 bp fragment of sequence centered on the ORF was passed to the primer picking software, PRIMER3 (available online for use at http://www-genome.wi.mit.edu/cgi-bin/primer/). A first additional sequence was commonly added to each ORF-unique 5' primer, and a second, different, additional sequence was commonly added to each ORF-unique 3' primer, to permit subsequent reamplification of the amplicon using a single set of "universal" 5' and 3' primers, thus immortalizing the amplicon. The addition of universal priming sequences also facilitates sequence verification, and can be used to add a cloning site should some ORFs be found to warrant further study.

The ORFs were then PCR amplified from genomic DNA, verified on agarose gels, and sequenced using the universal primers to validate the identity of the amplicon to be spotted in the microarray.

Primers were supplied by Operon Technologies (Alameda, CA). PCR amplification was performed by standard techniques using human genomic DNA (Clontech, Palo Alto, CA) as template. Each PCR product was verified by SYBR® green (Molecular Probes, Inc., Eugene, OR) staining of

agarose gels, with subsequent imaging by Fluorimager (Molecular Dynamics, Inc., Sunnyvale, CA). PCR amplification was classified as successful if a single band appeared.

The success rate for amplifying ORFs of interest directly from genomic DNA using PCR was approximately 75%. FIG. 5 graphs the distribution of predicted ORF (exon) length and distribution of amplified PCR products, with ORF length shown in red and PCR product length shown in blue (which may appear black in the figure). Although the range of ORF sizes is readily seen to extend to beyond 900 bp, the mean predicted exon size was only 229 bp, with a median size of 150 bp (n=9498). With an average amplicon size of 475 ± 25 bp, approximately 50% of the average PCR amplification product contained predicted coding region, with the remaining 50% of the amplicon containing either intron, intergenic sequence, or both.

Using a strategy predicated on amplifying about 500 bp, it was found that long exons had a higher PCR.

20 failure rate. To address this, the bioinformatics process was adjusted to amplify 1000, 1500 or 2000 bp fragments from exons larger than 500 bp. This improved the rate of successful amplification of exons exceeding 500 bp, constituting about 9.2% of the exons predicted by the gene finding algorithms.

Approximately 75% of the probes disposed on the array (90% of those that successfully PCR amplified) were sequence-verified by sequencing in both the forward and reverse direction using MegaBACE sequencer (Molecular Dynamics, Inc., Sunnyvale, CA), universal primers, and standard protocols.

Some genomic clones (BACs) yielded very poor PCR and sequencing results. The reasons for this are unclear, but may be related to the quality of early draft sequence or the inclusion of vector and host contamination in some

submitted sequence data.

Although the intronic and intergenic material flanking coding regions could theoretically interfere with hybridization during microarray experiments, subsequent empirical results demonstrated that differential expression ratios were not significantly affected by the presence of noncoding sequence. The variation in exon size was similarly found not to affect differential expression ratios significantly; however, variation in exon size was observed to affect the absolute signal intensity (data not shown).

The 350 MB of genomic DNA was, by the above-described process, reduced to 9750 discrete probes, which were spotted in duplicate onto glass slides using commercially available instrumentation (MicroArray GenII Spotter and/or MicroArray GenIII Spotter, Molecular Dynamics, Inc., Sunnyvale, CA). Each slide additionally included either 16 or 32 E. coli genes, the average hybridization signal of which was used as a measure of background biological noise.

Each of the probe sequences was BLASTed against the human EST data set, the NR data set, and SwissProt GenBank (May 7, 1999 release 2.0.9).

One third of the probe sequences (as amplified)

25 produced an exact match (BLAST Expect ("E") values less
than 1 e⁻¹⁰⁰) to either an EST (20% of sequences) or a known
mRNA (13% of sequences). A further 22% of the probe
sequences showed some homology to a known EST or mRNA
(BLAST E values from 1 e⁻⁵ to 1 e⁻⁹⁹). The remaining 45% of

30 the probe sequences showed no significant sequence homology
to any expressed, or potentially expressed, sequences
present in public databases.

All of the probe sequences (as amplified) were then analyzed for protein similarities with the SwissProt database using BLASTX, Gish et al., Nature Genet. 3:266

(1993). The predicted functional breakdowns of the 2/3 of probes identical or homologous to known sequences are presented in Table 1.

5 Table 1

Function	of Predic	ted ORFs As	Deduced From Comparative
Sequence	Analysis		
Total	V6 chip	V7 chip	Function Predicted from
			Comparative Sequence
			Analysis
211	96	115 .	Receptor
120	43	77	Zinc Finger
30	11	19	Homeobox
25	9	16	Transcription Factor
17	11	7	Transcription
118	57	61	Structural
95	39	56	Kinase
36	18	18	Phosphatase
83	31	52	Ribosomal
45	19	26	Transport
21	17	14	Growth Factor
17	12	5	Cytochrome
50	33	17	Channel

As can be seen, the two most common types of genes were transcription factors and receptors, making up 2.2% and 1.8% of the arrayed elements, respectively.

EXAMPLE 2

Gene Expression Measurements From Genome-Derived Single Exon Microarrays

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PCT/US01/00666 WO 01/57274

The two genome-derived single exon microarrays prepared according to Example 1 were hybridized in a series of simultaneous two-color fluorescence experiments to (1) 5 Cy3-labeled cDNA synthesized from message drawn individually from each of brain, heart, liver, fetal liver, placenta, lung, bone marrow, HeLa, BT 474, or HBL 100 cells, and (2) Cy5-labeled cDNA prepared from message pooled from all ten tissues and cell types, as a control in 10 each of the measurements. Hybridization and scanning were carried out using standard protocols and Molecular Dynamics equipment.

Briefly, mRNA samples were bought from commercial sources (Clontech, Palo Alto, CA and Amersham Pharmacia 15 Biotech (APB)). Cy3-dCTP and Cy5-dCTP (both from APB) were incorporated during separate reverse transcriptions of 1 µg of polyA+ mRNA performed using 1 µg oligo(dT)12-18 primer and 2 µg random 9mer primers as follows. After heating to 70°C, the RNA:primer mixture was snap cooled on ice. After 20 snap cooling on ice, added to the RNA to the stated final concentration was: 1X Superscript II buffer, 0.01 M DTT, 100μM dATP, 100 μM dGTP, 100 μM dTTP, 50 μM dCTP, 50 μM Cy3-dCTP or Cy5-dCTP 50 µM, and 200 U Superscript II The reaction was incubated for 2 hours at 42°C. 25 After 2 hours, the first strand cDNA was isolated by adding 1 U Ribonuclease H, and incubating for 30 minutes at 37°C. The reaction was then purified using a Qiagen PCR cleanup column, increasing the number of ethanol washes to 5. Probe was eluted using 10 mM Tris pH 8.5.

Using a spectrophotometer, probes were measured for dye incorporation. Volumes of both Cy3 and Cy5 cDNA corresponding to 50 pmoles of each dye were then dried in a Speedvac, resuspended in 30 µl hybridization solution containing 50% formamide, 5X SSC, 0.2 $\mu g/\mu l$ poly(dA), 0.2 35 $\mu g/\mu l$ human $c_0 t l$ DNA, and 0.5 % SDS.

Hybridizations were carried out under a coverslip, with the array placed in a humid oven at 42°C overnight. Before scanning, slides were washed in 1X SSC, 0.2% SDS at 55° C for 5 minutes, followed by 0.1X SSC, 0.2%5 SDS, at 55°C for 20 minutes. Slides were briefly dipped in water and dried thoroughly under a gentle stream of nitrogen.

Slides were scanned using a Molecular Dynamics Gen3 scanner, as described. Schena (ed.), Microarray 10 Biochip: Tools and Technology, Eaton Publishing Company/BioTechniques Books Division (2000) (ISBN: 1881299376).

Although the use of pooled cDNA as a reference permitted the survey of a large number of tissues, it 15 attenuates the measurement of relative gene expression, since every highly expressed gene in the tissue/cell typespecific fluorescence channel will be present to a level of at least 10% in the control channel. Because of this fact, both signal and expression ratios (the latter hereinafter, 20 "expression" or "relative expression") for each probe were normalized using the average ratio or average signal, respectively, as measured across the whole slide.

Data were accepted for further analysis only when signal was at least three times greater than biological 25 noise, the latter defined by the average signal produced by the E. coli control genes.

The relative expression signal for these probes was then plotted as function of tissue or cell type, and is presented in FIG. 6.

FIG. 6 shows the distribution of expression across a panel of ten tissues. The graph shows the number of sequence-verified products that were either not expressed ("0"), expressed in one or more but not all tested tissues ("1" - "9"), and expressed in all tissues 35 tested ("10").

Of 9999 arrayed elements on the two microarrays (including positive and negative controls and "failed" products), 2353 (51%) were expressed in at least one tissue or cell type. Of the gene elements showing significant signal — where expression was scored as "significant" if the normalized Cy3 signal was greater than 1, representing signal 5-fold over biological noise (0.2) — 39% (991) were expressed in all 10 tissues. The next most common class (15%) consisted of gene elements expressed in only a single tissue.

The genes expressed in a single tissue were further analyzed, and the results of the analyses are compiled in FIG. 7.

FIG. 7A is a matrix presenting the expression of 15 all verified sequences that showed expression greater than 3 in at least one tissue. Each clone is represented by a column in the matrix. Each of the 10 tissues assayed is represented by a separate row in the matrix, and relative expression of a clone in that tissue is indicated at the 20 respective node by intensity of green shading, with the intensity legend shown in panel B. The top row of the matrix ("EST Hit") contains "bioinformatic" rather than "physical" expression data - that is, presents the results returned by query of EST, NR and SwissProt databases using 25 the probe sequence. The legend for "bioinformatic expression" (i.e., degree of homology returned) is presented in panel C. Briefly, white is known, black is novel, with gray depicting nonidentical with significant homology (white: E values < 1e-100; gray: E values from 1e-30 05 to 1e-99; black: E values > 1e-05).

As FIG. 7 readily shows, heart and brain were demonstrated to have the greatest numbers of genes that were shown to be uniquely expressed in the respective tissue. In brain, 200 uniquely expressed genes were identified; in heart, 150. The remaining tissues gave the

following figures for uniquely expressed genes: liver, 100; lung, 70; fetal liver, 150; bone marrow, 75; placenta, 100; HeLa, 50; HBL, 100; and BT474, 50.

It was further observed that there were many more

"novel" genes among those that were up-regulated in only
one tissue, as compared with those that were down-regulated
in only one tissue. In fact, it was found that ORFs whose
expression was measurable in only a single of the tested
tissues were represented in sequencing databases at a rate
of only 11%, whereas 36% of the ORFs whose expression was
measurable in 9 of the tissues were present in public
databases. As for those ORFs expressed in all ten tissues,
fully 45% were present in existing expressed sequence
databases. These results are not unexpected, since genes
expressed in a greater number of tissues have a higher
likelihood of being, and thus of having been, discovered by
EST approaches.

Comparison of Signal from Known and Unknown Genes

The normalized signal of the genes found to have high homology to genes present in the GenBank human EST database were compared to the normalized signal of those genes not found in the GenBank human EST database. The data are shown in FIG. 8.

FIG. 8 shows the normalized Cy3 signal intensity for all sequence-verified products with a BLAST Expect ("E") value of greater than 1e-30 (designated "unknown") upon query of existing EST, NR and SwissProt databases, and shows in blue the normalized Cy3 signal intensity for all sequence-verified products with a BLAST Expect value of less than 1e-30 ("known"). Note that biological background noise has an averaged normalized Cy3 signal intensity of 0.2.

As expected, the most highly expressed of the 35 ORFs were "known" genes. This is not surprising, since

very high signal intensity correlates with very commonlyexpressed genes, which have a higher likelihood of being found by EST sequence.

However, a significant point is that a large

5 number of even the high expressers were "unknown". Since
the genomic approach used to identify genes and to confirm
their expression does not bias exons toward either the 3'
or 5' end of a gene, many of these high expression genes
will not have been detected in an end-sequenced cDNA

10 library.

The significant point is that presence of the gene in an EST database is not a prerequisite for incorporation into a genome-derived microarray, and further, that arraying such "unknown" exons can help to assign function to as-yet undiscovered genes.

Verification of Gene Expression

To ascertain the validity of the approach described above to identify genes from raw genomic

20 sequence, expression of two of the probes was assayed using reverse transcriptase polymerase chain reaction (RT PCR) and northern blot analysis.

Two microarray probes were selected on the basis of exon size, prior sequencing success, and tissue-specific gene expression patterns as measured by the microarray experiments. The primers originally used to amplify the two respective ORFs from genomic DNA were used in RT PCR against a panel of tissue-specific cDNAs (Rapid-Scan gene expression panel 24 human cDNAs) (OriGene Technologies, Inc., Rockville, MD).

Sequence AL079300_1 was shown by microarray hybridization to be present in cardiac tissue, and sequence AL031734_1 was shown by microarray experiment to be present in placental tissue (data not shown). RT-PCR on these two sequences confirmed the tissue-specific gene expression as

measured by microarrays, as ascertained by the presence of a correctly sized PCR product from the respective tissue type cDNAs.

Clearly, all microarray results cannot, and
indeed should not, be confirmed by independent assay
methods, or the high throughput, highly parallel advantages
of microarray hybridization assays will be lost. However,
in addition to the two RT-PCR results presented above, the
observation that 1/3 of the arrayed genes exist in
expression databases provides powerful confirmation of the
power of our methodology — which combines bioinformatic
prediction with expression confirmation using genome—
derived single exon microarrays — to identify novel genes
from raw genomic data.

To verify that the approach further provides correct characterization of the expression patterns of the identified genes, a detailed analysis was performed of the microarrayed sequences that showed high signal in brain.

For this latter analysis, sequences that showed
20 high (normalized) signal in brain, but which showed very
low (normalized) signal (less than 0.5, determined to be
biological noise) in all other tissues, were further
studied. There were 82 sequences that fit these criteria,
approximately 2% of the arrayed elements. The 10 sequences
25 showing the highest signal in brain in microarray
hybridizations are detailed in Table 2, along with assigned
function, if known or reasonably predicted.

Table 2

15

Function of the Most Highly Expressed Genes Expressed Only in Brain

Microarray	Normal	Expressi	Homology	Gene Function
Sequence	ized	on Ratio	to EST	as described by
Name .	Signal		present	GenBank
			in	
			GenBank	
AP000217-1	5.2	+7.7	High	S-100 protein,
				b-chain, Ca ²⁺
				binding protein
				expressed in
				central nervous
				system
AP000047-1	2.3		High	Unknown
	ı			Function
AC006548-9	1.7		High	Similar to
				mouse membrane
				glyco-protein
				M6, expressed
				in central
				nervous system
AC007245-5	1.5		High	Similar to
				amphiphysin, æ
				synaptic
				vesicle-
				associated
				protein. Ref 21
L44140-4	1.2	+2.0	High	Endothelial
·				actin-binding
				protein found
				in nonmuscle
				filamin

AC004689-9	1.2	+3.5	High	Protein
				Phosphatase
				PP2A, neuronal/
		:		downregulates
				activated
				protein kinases
AL031657-1	1.2	+3.0	High	Unknown
				function/
				Contains the
•				anhyrin motif,
		ĺ		a common
	<u>.</u>	[protein
		:		sequence motif
AC009266-2	1.1	+3.7	Low	Low homology to
	,			the
				Synaptotagmin I
				protein in
				rat/present at
				low levels
				throughout rat
:				brain
AP000086-1	1.0	+2.7	Low	Unknown, very
-				poor homology
				to collagen
AC004689-3	1.0		High	Protein
				Phosphatase
				PP2A, neuronal/
				downregulates
				activated
·				protein kinases

Of the ten sequences studied by these latter confirmatory approaches, eight were previously known. Of these eight, six had previously been reported to be important in the central nervous system or brain. The exon

giving the highest signal (AP00217-1) was found to be the gene encoding an S100B Ca²⁺ binding protein, reported in the literature to be highly and uniquely expressed in the central nervous system. Heizmann, Neurochem. Res. 9:1097 (1997).

A number of the brain-specific probe sequences (including AC006548-9, AC009266-2) did not have homology to any known human cDNAs in GenBank but did show homology to rat and mouse cDNAs. Sequences AC004689-9 and AC004689-3

10 were both found to be phosphatases present in neurons (Millward et al., Trends Biochem. Sci. 24(5):186-191 (1999)). Two microarray sequences, AP000047-1 and AP000086-1 have unknown function, with AP000086-1 being absent from GenBank. Functionality can now be narrowed down to a role in the central nervous system for both of these genes, showing the power of designing microarrays in this fashion.

Next, the function of the chip sequences with the highest (normalized) signal intensity in brain, regardless of expression in other tissues, was assessed. In this latter analysis, we found expression of many more common genes, since the sequences were not limited to those expressed only in brain. For example, looking at the 20 highest signal intensity spots in brain, 4 were similar to tubulin (AC00807905; AF146191-2; AC007664-4; AF14191-2), 2 were similar to actin (AL035701-2; AL034402-1), and 6 were found to be homologous to glyceraldehyde-3-phosphate dehydrogenase (GAPDH) (AL035604-1; Z86090-1; AC006064-L, AC006064-K; AC035604-3; AC006064-L). These genes are often used as controls or housekeeping genes in microarray experiments of all types.

Other interesting genes highly expressed in brain were a ferritin heavy chain protein, which is reported in the literature to be found in brain and liver (Joshi et al., J. Neurol. Sci. 134(Suppl):52-56 (1995)), a result

duplicated with the array. Other highly expressed chip sequences included a translation elongation factor 1D (AC007564-4), a DEAD-box homolog (AL023804-4), and a Y-chromosome RNA-binding motif (Chai et al., Genomics 49(2):283-89 (1998)) (AC007320-3). A low homology analog (AP00123-1/2) to a gene, DSCR1, thought to be involved in trisomy 21 (Down's syndrome), showed high expression in both brain and heart, in agreement with the literature (Fuentes et al., Mol. Genet. 4(10):1935-44 (1995)).

As a further validation of the approach, we selected the BAC AC006064 to be included on the array. This BAC was known to contain the GAPDH gene, and thus could be used as a control for the ORF selection process. The gene finding and exon selection algorithms resulted in choosing 25 exons from BAC AC006064 for spotting onto the array, of which four were drawn from the GAPDH gene. Table 3 shows the comparison of the average expression ratio for the 4 exons from BAC006064 compared with the average expression ratio for 5 different dilutions of a commercially available GAPDH cDNA (Clontech).

Table 3

Comparis	son of Expression Ra	atio, for each
tissue, of GAPDH		
	AC006064 (n = 4)	Control (n = 5)
Bone Marrow	-1.81 ± 0.11	-1.85 ± 0.08
Brain	-1.41 ± 0.11	-1.17 ± 0.05
BT474	1.85 ± 0.09	1.66 ± 0.12
Fetal Liver	-1.62 ± 0.07	-1.41 ± 0.05
HBL100	1.32 ± 0.05	2.64 ± 0.12
Heart	1.16 ± 0.09	1.56 ± 0.10
HeLa	1.11 ±0.06	1.30 ± 0.15
Liver	-1.62 ± 0.22	-2.07 ±

Lung	-4.95 ± 0.93	-3.75 ± 0.21
Placenta	-3.56 ± 0.25	-3.52 ± 0.43

Each tissue shows excellent agreement between the experimentally chosen exons and the control, again

5 demonstrating the validity of the present exon mining approach. In addition, the data also show the variability of expression of GAPDH within tissues, calling into question its classification as a housekeeping gene and utility as a housekeeping control in microarray experiments.

EXAMPLE 3

Representation of Sequence and Expression Data as a "Mondrian"

15

For each genomic clone processed for microarray as above-described, a plethora of information was accumulated, including full clone sequence, probe sequence within the clone, results of each of the three gene finding programs, EST information associated with the probe sequences, and microarray signal and expression for multiple tissues, challenging our ability to display the information.

Accordingly, we devised a new tool for visual display of the sequence with its attendant annotation which, in deference to its visual similarity to the paintings of Piet Mondrian, is hereinafter termed a "Mondrian". FIGS. 3 and 4 present the key to the information presented on a Mondrian.

FIG. 9 presents a Mondrian of BAC AC008172 (bases 25,000 to 130,000 shown), containing the carbamyl phosphate synthetase gene (AF154830.1). Purple background within the region shown as field 81 in FIG. 3 indicates all 37 known

exons for this gene.

As can be seen, GRAIL II successfully identified 27 of the known exons (73%), GENEFINDER successfully identified 37 of the known exons (100%), while DICTION identified 7 of the known exons (19%).

Seven of the predicted exons were selected for physical assay, of which 5 successfully amplified by PCR and were sequenced. These five exons were all found to be from the same gene, the carbamyl phosphate synthetase gene (AF154830.1).

The five exons were arrayed, and gene expression measured across 10 tissues. As is readily seen in the Mondrian, the five chip sequences on the array show identical expression patterns, elegantly demonstrating the reproducibility of the system.

FIG. 10 is a Mondrian of BAC AL049839. We selected 12 exons from this BAC, of which 10 successfully sequenced, which were found to form between 5 and 6 genes. Interestingly, 4 of the genes on this BAC are protease 20 inhibitors. Again, these data elegantly show that exons selected from the same gene show the same expression patterns, depicted below the red line. From this figure, it is clear that our ability to find known genes is very good. A novel gene is also found from 86.6 kb to 88.6 kb, 25 upon which all the exon finding programs agree. We are confident we have two exons from a single gene since they show the same expression patterns and the exons are proximal to each other. Backgrounds in the following colors indicate a known gene (top to bottom): 30 red = kallistatin protease inhibitor (P29622); purple = plasma serine protease inhibitor (P05154); turquoise = α 1 anti-chymotrypsin (P01011); mauve = 40S ribosomal protein (P08865). Note that chip sequence 8 and 12 did not sequence verify.

EXAMPLE 4

Genome-Derived Single Exon Probes Useful For Measuring Human Gene Expression

5

The protocols set forth in Examples 1 and 2, supra, were applied to additional human genomic sequence as it became newly available in GenBank to identify unique exons in the human genome that could be shown to be expressed at significant levels in heart tissue.

These unique exons are within longer probe sequences. Each probe was completely sequenced on both strands prior to its use on a genome-derived single exon microarray; sequencing confirms the exact chemical

15 structure of each probe. An added benefit of sequencing is that it placed us in possession of a set of single base-incremented fragments of the sequenced nucleic acid, starting from the sequencing primer 3' OH. (Since the single exon probes were first obtained by PCR amplification from genomic DNA, we were of course additionally in possession of an even larger set of single base incremented fragments of each of the 9,980 single exon probes, each fragment corresponding to an extension product from one of the two amplification primers.)

25 The structures of the 9,980 unique single exon probes are clearly presented in the Sequence Listing as SEQ ID Nos.: 1 - 9,980. The 16 nt 5' primer sequence and 16 nt 3' primer sequence present on the amplicon are not included in the sequence listing. The sequences of the exons present within each of these probes is presented in the Sequence Listing as SEQ ID Nos.: 9,981 - 19,771, respectively. It will be noted that some amplicons have more than one exon, some exons are contained in more than one amplicon.

As detailed in Example 2, expression was

demonstrated by disposing the amplicons as single exon probes on nucleic acid microarrays and then performing two-color fluorescent hybridization analysis; significant expression is based on a statistical confidence that the signal is significantly greater than negative biological control spots. The negative biological control is formed from spotted DNA sequences from a different species. Here, 32 sequences from E.Coli were spotted in duplicate to give a total of 64 spots.

10 For each hybridisation (each slide, each colour) the median value of the signal from all of the spots is determined. The normalised signal value is the arithmetic mean of the signal from duplicate spots divided by the population median.

Control spots are eliminated if there is more that a five-fold difference between each one of the duplicate spots raw signals.

15

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The median of the signal from the remaining control spots is calculated and all subsequent calculations are done with normalised signals.

Control spots having a signal of greater than median + 2.4 (the value 2.4 is roughly 12 times the observed standard deviation of control spot populations) are eliminated. Spots with such high signals are considered to be "outliers".

The mean and standard deviation of the modified control spot populations are calculated.

The mean + 3x the standard deviation (mean + (3*SD)) is used as the signal threshold qualifier for that particular hybridisation. Thus, individual thresholds are determined for each channel and each hybridisation.

This means that, assuming that the data is distributed normally, there is a 99% confidence that any signal exceeding the threshold is significant.

The probes and their expression data are

presented in Table 4, set forth respectively in Example 5.

Example 5 presents the subset of probes that is significantly expressed in the human heart and thus presents the subset of probes that was recognized to be useful for measuring expression of their cognate genes in human heart tissue.

The sequence of each of the exon probes identified by SEQ ID NOS.: 9,981 - 19,771 was individually used as a BLAST (or, for SWISSPROT, BLASTX) query to identify the most similar sequence in each of dbEST, SwissProt (BLASTX), and NR divisions of GenBank. Because the query sequences are themselves derived from genomic sequence in GenBank, only nongenomic hits from NR were scored.

15

The smallest in value of the BLAST (or BLASTX) expect ("E") scores for each query sequence across the three database divisions was used as a measure of the "expression novelty" of the probe's ORF. Table 4 is sorted in descending order based on this measure, reported as "Most Similar (top) Hit BLAST E Value". Those sequences for which no "Hit E Value" is listed are those exons which were found to have no similar sequences.

As sorted, Table 4 thus lists its respective probes (by "AMPLICON SEQ ID NO.:" and additionally by the SEQ ID NO.: of the exon contained within the probe: "EXON SEQ ID NO.:") from least similar to sequences known to be expressed (i.e., highest BLAST E value), at the beginning of the table, to most similar to sequences known to be expressed (i.e., lowest BLAST E value), at the bottom of the table.

Table 4 further provides, for each listed probe, the accession number of the database sequence that yielded the "Most Similar (top) Hit BLAST E Value", along with the name of the database in which the database sequence is found ("Top Hit Database Source").

Table 4 further provides SEQ ID NOS. corresponding to the predicted amino acid sequences where they have been determined for the probe and exon nucleotide sequences. These are set out as PEPTIDE SEQ ID NOS.:. The 5 peptide sequences for a given exon are predicted as follows: Since each chip exon is a consensus sequence drawn from predictions from various exon finding programs (i.e. Grail, GeneFinder and GenScan), the multiple initial ORFs are first determined in a uniform way according to each 10 prediction. In particular, the reading frame for predicting the first amino acid in the peptide sequence always starts with the first base of any codon and ends with the last base of non-termination codon. Next, for each strand of the exon, initial ORFs are merged into one or more final ORFs 15 in an exhaustive process based on the following criteria: 1) the merging ORFs must be overlapping, and 2) the merging ORFs must be in the same frame.

The Sequence Listing, which is a superset of all of the data presented in Table 4, further includes, for 20 each probe, the most similar hit, with accession number and BLAST E value, from the each of the three queried databases.

Table 4 further lists, for each probe, a portion of the descriptor for the top hit ("Top Hit Descriptor") as 25 provided in the sequence database. For those ORFs that are similar in sequence, but nonidentical to known sequences (e.g., those with BLAST E values between about 1e-05 and 1e-100), the descriptor reveals the likely function of the protein encoded by the probe's ORF.

Using BLAST E value cutoffs of 1e-05 (i.e., 1 x 10^{-5}) and le-100 (i.e., 1 x 10^{-100}) as evidence of similarity to sequences known to be expressed is of course arbitrary: in Example 2, supra, a BLAST E value of 1e-30 was used as the boundary when only two classes were to be defined for 35 analysis (unknown, >1e-30; known <1e-30) (see also FIG. 8).

Furthermore, even when the "Most Similar (Top) Hit BLAST E Value" is low, e.g., less than about 1e-100 — which is probative evidence that the query sequence has previously been shown to be expressed — the top hit is highly unlikely exactly to match the probe sequence.

First, such expression entries typically will not have the intronic and/or intergenic sequence present within the single exon probes listed in the Table. Second, even the ORF itself is unlikely in such cases to be present identically in the databases, since most of the EST and mRNA clones in existing databases include multiple exons, without any indication of the location of exon boundaries.

As noted, the data presented in Table 4 represent a proper subset of the data present within the attached sequence listing. For each amplicon probe (SEQ ID NOs.: 1 - 9,980) and probe exon (SEQ ID NOs.: 9,981 - 19,771, respectively), the sequence listing further provides, through iterated annotation fields <220> and <223>:

- . . (a) the accession number of the BAC from which

 the sequence was derived ("MAP TO"), thus providing a link

 to the chromosomal map location and other information about

 the genomic milieu of the probe sequence;
- (b) the most similar sequence provided by BLAST query of the EST database, with accession number and BLAST 25 E value for the "hit";
 - (c) the most similar sequence provided by BLAST query of the GenBank NR database, with accession number and BLAST E value for the "hit"; and
- (d) the most similar sequence provided by BLASTX 30 query of the SWISSPROT database, with accession number and BLAST E value for the "hit".

EXAMPLE 5

35 Genome-Derived Single Exon Probes Useful For Measuring

Expression of Genes in Human Heart

<u>Table 4</u> (413 pages) presents expression, homology, and functional information for the genome-derived single exon probes that are expressed significantly in human heart.

Page 1 of 413 Table 4 Single Exon Probes Expressed in Heart

	Top Hit Descriptor						•																												
201 1 1000	Top Hit Database Source																																		
Sign of	Top Hit Acesslon No.																																		
	Most Similar (Top) Hit BLAST E Value																																		
	Expression Signal	4.41	17.08	2.14	7.97	1.87	4.97	1.01	0.95	7.45	96.0	3.03	2.62	2.34	3.7	1.52	8.97	. 0.87	66.0	1.71	. 5.94	0.89	0.89	1.03	1.76	0.78	5.08	2.07	1.92	1.92	9	5.89	4.9	29	1.6
	ORF SEQ ID NO:	20209	20647		21039	21361			21485		21620		ĺ						23309				23859		24402		24617	24629		24835					
	Exon SEQ ID NO:	10386	10797	10947	11188		11523		11616	11622	11744	11830	ı	12118	<u> </u>			13428	13521	14002	14064	14084	14084	14141		14663	14851	14863	15140	15140	15248		15248		19451
	Probe SEQ ID NO:	442	871	1029	1280	1597	1619	1694	1715	1721	1848	1935	2119	2233	3149	3403	3471	3513	3607	4102	4164	4184	4184	4242	4730	4779	4976	4988	5217	5217	5328	5436	5483	5509	5668

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Table 4
Single Exon Probes Expressed in Heart

Г						Γ-		Γ-	П					_						_		$\overline{}$			_	г			
	Top HIt Descriptor																		Homo sapiens LSS gene, partial, exons 15, 16, 17 and 18	Haemophilus influenzae Rd section 31 of 163 of the complete genome	Mus musculus Naip3 gene, exon 1; neuronal apoptosis inhibitory protein 1 (Naip1) and general transcription factor IIH polypeptide 2 (Gtf2h2) genes, complete cds	Mus musculus Naip3 gene, exon 1; neuronal apoptosis inhibitory protein 1 (Naip1) and general transcription factor IIH polypeptide 2 (Gtf2h2) genes, complete cds	Dengue virus type 3 membrane protein (priM/M)/envelope glycoprotein (E) polyprotein mRNA, partial cds	Dengue virus type 3 membrane protein (prM/M)/envelope glycoprotein (E) polyprotein mRNA, partial cds	Mus musculus AT3 gene for antithrombin, complete cds	IMMEDIATE-EARLY PROTEIN 1 (IE1) (IMMEDIATE-EARLY PHOSPHOPROTEIN PP89)	Leuciscus cephalus orientalis cytochrome b (cyt.b) gene, partial cds; mitochondrial gene for mitochondrial product	Leuciscus cephalus orientalis cytochrome b (cyt.b) gene, partial ods; mitochondrial gene for mitochondrial product.	601651038R1 NIH_MGC_81 Homo sapiens cDNA clone IMAGE:3934592 3'
	Top Hit Database Source																		ΙN	F	F	뉟	NT	Ę	L.	SWISSPROT	N	ΤN	EST HUMAN
	Top Hit Acession No.																		AJ239028.1	U32716.1	AF242432.1	AF242432.1	L11433.1	L11433.1	AB043785.1	P11210	AF095809.1	AF095509.1	BE971806.1
	Most Similar (Top) Hit BLAST E Value																		9.9E+00	9.8E+00	9.6E+00	9.6E+00	9.4E+00	9.4E+00 [9.3E+00	9.1E+00	9.1E+00	
	Expression Signal	1.41	1.84	1.61	1.61	1.29	1.3	3.58	1.28		2.31	2.84	2.91	1.73	1.73	2.19	1.62	2.11	13.31	1.54	1.32	1.32	2.72	2.72	5.91	2.97	2.07	2.07	5.51
	ORF SEQ ID NO:	25700		26404	28405	26769	27012	27569	27758		28366		28840	28150	28151			25230	25701	26745	28046	28047	22389	22390		27271	24932	24933	25687
	Exon SEQ ID NO: .	15599	15769	16244	16244	16577	16820	17364	17534	17861	19472	18268	18555	17906	17808	18608	19094	19280	15600	16549	17805	17805	12499	12499		17083	15163	15163	l i
	Probe SEQ ID NO:	2690	5863	6382	6382	6697	6942	7494	7684	8070	8219	8390	8868	8757	8757	8792	9464	9760	5691	6999	7955	7955	2631	2631	2893	7206	5239	. 6239	5678

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Table 4
Single Exon Probes Expressed in Heart

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acesslan No.	Top Hit Database Source	Top Hit Descriptor
5848	15754	25871	1.71		AB019788.1	TN	Cynops pyrrhogaster CpTbx3 premature mRNA, partial cds
5848	15754	25872	1.71	8.7E+00	AB019788.1	TN.	Oynops pyrrhogaster OpTbx3 premature mRNA, partial cds
433	乚	20199	1.76	8.4E+00	5031804 NT	IN	Homo saplens insulin receptor substrate 1 (IRS₄) mRNA
7439	16452	26642	3.68		AJ131719.1	LN	Zea mays mRNA for legumain-like protease (sea2a)
8509	18381		2.31	8.0E+00	P41820	SWISSPROT	BREFELDIN A RESISTANCE PROTEIN
6346	16209	,	2.07	7.5E+00	35.1	TN	Thermoplasma acidophilum complete genome; segment 3/5
8989	16747	26940	1.53			SWISSPROT	THROMBOSPONDIN 1 PRECURSOR
6868	16747	26941	1.53	7.5E+00	P35441	SWISSPROT	THROMBOSPONDIN 1 PRECURSOR
5544	15460		2.58	7.4E+00	BF700517.1	EST_HUMAN	602128876F1 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:4285506 5'
7085	16962		3.83	7.4E+00	P04929	SWISSPROT	HİSTIDINE-RICH GLYCOPROTEIN PRECURSOR
7085	16962		3.83		P04929	SWISSPROT	HISTIDINE-RICH GLYCOPROTEIN PRECURSOR
2947	12874	22671	4.2	7.2E+00	L12051.1	IN	Lycopersicon esculentum Mill. GTPase (SAR2) mRNA, complete cds
2947	12874	22672	4.2	7.2E+00	L12051.1	NT	Lycopersicon esculentum Mill. GTPase (SAR2)/mRNA, complete cds
6239	16105		1.3	7.1E+00	P28166	SWISSPROT	ZINC-FINGER PROTEIN 1 (ZINC-FINGER HOMEODOMAIN PROTEIN 1)
6238	16105	26255	1.3	7.1E+00	P28166	SWISSPROT	ZINC-FINGER PROTEIN 1 (ZINC-FINGER HOMEODOMAIN PROTEIN 1)
7535	17386		8,48	7.1E+00	AL161595.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 91
8690	18577	28860	3.26	7.1E+00	P05850	SWISSPROT	HYPOTHETICAL 17.3 KDA PROTEIN IN MRDA-PHPB INTERGENIC REGION
7735	17685	27809	2.98	7.0E+00	P48610	SWISSPROT	ARGININE KINASE (AK)
8575	18443	28712	1.85	7.0E+00	022469	SWISSPROT	WD-40 REPEAT PROTEIN MSI3
6818	16697	26889	2.72	6.9E+00	P35679	SWISSPROT	60S RIBOSOMAL PROTEIN L4 (L2)
7925	17775		1.3		P44834	SWISSPROT	DNA MISMATCH REPAIR PROTEIN MUTS
6614	16494	26680	1.45	00+38'9	W03412.1	EST_HUMAN	za07c11.r1 Soares melanocyte 2NbHM Homo saplens cDNA clone IMAGE:291860 5'
6614	18494	26681	1.45	0.8E+00	W03412.1	EST_HUMAN	za07c11.r1 Soares melanocyte 2NbHM Homo sapiens cDNA clone IMAGE:291860 5'
	L		,				OUTER CAPSID PROTEIN VP4 (HEMAGGLUTININ) (OUTER LAYER PROTEIN VP4) [CONTAINS:
//2/	┙			6.8E+00		SWISSPROI	OUTEN CATE OF THE STATE OF THE
7883	17713			6.8E+00	Q035/0	SWISSPROI	TITOTHETICAL 197.0 KDA TROTEIN CSSCIOSIN CHROMOSOME III TIBIDATATE KINASE 71KV (TIBIDINE MONOBHOSOHATE KINASEV (TIMD KINASEV
7706	L	27001	2,5	_	G92E07	SWISSPACE	INDICATE KINDSE (1K) (I IRIDINE MONOPHOSPHATE KINDSE) (I IMP KINDSE)
26,70	L				240300	TOGGSSIWS	PROBABLE CATION-TRANSPORTING ATPASE CACA 050
	L			0.17.0	2,0000	FOOD SOLVE	ENV DOI YEDOTEN CONTAINS: COAT BEOTEIN GES- COAT BEOTEIN CEPTRE
087/		Ì		0.00+100		DA POST WO	
7616	17467	27686	1.44	6.2E+00	AY010901.1	Ę	Schizophyllum commune unknown mRNA
				L			Mus musculus mixed lineage kinase 3 (MiK3) and two pore domain K+ channel subunit (Konk6) genes,
5928	ı	25956		5.9E+00	AF165142.1	L !	complete cds
3479	13395		0.81	5.8E+00	7861557 NT	Į.	Homo sapiens DESC1 protein (DESC1), mKNA

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Top Hit Descriptor	LYCOPENE BETA CYCLASE	PNEUMOLYSIN (THIOL-ACTIVATED CYTOLYSIN)	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 67	VITELLOGENIN PRECURSOR (VTG) [CONTAINS: LIPOVITELLIN LV-1N; LIPOVITELLIN LV-1C;	[LIPOVITELLIN LV-2]	RHODOPSIN	RHODOPSIN	Bovine immunodeficiency-like virus surface envelope gene, 5' end of cds	DNA POLYMERASE GAMMA (MITOCHONDRIAL DNA POLYMERASE CATALYTIC SUBUNIT)	PROBABLE ANTIBACTERIAL PEPTIDE POLYPROTEIN PRECURSOR	COLICIN N IMMUNITY PROTEIN (MICROCININ IMMUNITY PROTEIN)	Canis familiaris skeletal muscle chloride channel CIC-1 (CLCN1) mRNA, complete cds	Mycobacterium tuberculosis H37Rv complete genome; segment 103/162	Eunice australis histone H3 (H3) gene, partial cds	PM0-BT0547-310100-002-b04 BT0547 Homo sapiens cDNA	601875654F1 NIH_MGC_55 Homo septens cDNA clone IMAGE:4099716 5'	601875654F1 NIH_MGC_55 Homo sapiens cDINA clone IMAGE:4099716 5	Homo sapiens chromosome 21 segment HS21C080	Methanococcus Jannaschil section 111 of 150 of the complete genome	Archaeoglobus fulgidus section 63 of 172 of the complete genome	[602072585F1 NCI_CGAP_Bm67 Homo saplens cDNA clone IMAGE:4215284 5'	602072585F1 NCL_CGAP_Bm67 Homo saplens cDNA clone IMAGE:4215284 5'	Murine I gene for MHC class II(Ia) associated invariant chain	Plasmodium falciparum R29R+var1 gene, exon it	Homo sapiens glutathione S-transferase theta 2 (GSTT2) and glutathione S-transferase theta 1 (GSTT1)	genes, complete cds	MICROSOMAL DIPEPTIDASE PRECURSOR (MDP) (DEHYDROPEPTIDASE-1) (RENAL DIPEPTIDASE) (RDP)	EXTENSIN DRED (DROP (CELL WAIT HYDROXYBBO) INE BICH CLYCODBOTEIN)	EXTENSIVE TRANSPORT OF THE WALL THE PROPERTY OF THE PROPERTY O	EXTENSIN PRECORSOR (CELL WALL INTOROXIPECHINE-RICH GLYCOPROTEIN)	wf67g03.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2360692.31	YY1 PROTEIN PRECURSOR	GENE 68 PROTEIN	GENE 68 PROTEIN
Top Hit Database Source	SWISSPROT	SWISSPROT	Ā		SWISSPROT	SWISSPROT	SWISSPROT	ΙN	SWISSPROT	SWISSPROT	SWISSPROT	NT	TN	ΝŢ	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	INT	NT	EST_HUMAN	EST_HUMAN	NT	NT		NT	TORGSSIWS	TOUCOUNT	SWISSPROI	SWISSPROI	EST HUMAN	SWISSPROT	SWISSPROT	SWISSPROT
Top Hit Acession No.	Q55278	P11890	AL161571.2		Q91062	Q17094	Q17094	L43126.1	P54098	Q27905	P09182	AF162445.2	Z83860.1	AF186265.1	AW750067.1	BF240552.1	BF240552.1	AL163280.2	U67569.1	AE001044.1	BF530893.1	BF530893.1	X13414.1	Y13402.1		AF240786.1	P16444	27000	713805	P13983	A1809013.1	023810	P28964	P28964
Most Similar (Тор) Hit BLAST E Value	5.6E+00	5.5E+00	6.5E+00		5.4E+00	5.4E+00	5.4E+00	5.3E+00	8.3E+00	5.3E+00	5.1E+00			4.8E+00	4.8E+00	4.7E+00	4.7E+00	4.7E+00		4.5E+00	4.4E+00	4.4E+00	4.4E+00	4.3E+00		4.3E+00	4 25+00		4.ZE+00				4.1E+00	4.1E+00
Expression Signal	2.44	2.65	1.94		1.62	1.44	1.44	1.54	4.04	3.21	1.21	3.06	10.53	9.43	5.01	2.03	1.92	4.01	1.48	1.99	0.96	96.0	1.55	2.01		7.49	200		1.5/	1.57	5.45	7.65	3.31	3.31
ORF SEQ ID NO:		28165				27849	27850	24354		28003	27730	28050	28749			20071			26617	28986		22724		26420		28368		1	1					26619
Exon SEQ ID NO:	17919	17918	18773	L	16473	17619	17619	14581	16590	18708	17505	17809	18477	13879	16847	10251			16434	18693	12931	12931	12671	16259		18116	15306	1		- 1			16435	16435
Probe SEQ ID NO:	8770	8769	8967		6583	7769	7769	4675	6710	6688	7655	7959	8610	3972	6970	286	287	3236	8576	8881	3003	3003	5764	8338		8235	K3.8.7		200	0909	7182	6517	6577	6577

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Probe SEC ID 10 NO. Expression ID NO. Most Similar No. Top Hit No. Top Hit No. Top Hit No. Top Hit No. Top Hit No. Top Hit No. Top Hit Sucrate Top Hit Descriptor 8167.1 (ASP) 7375 2.897 (ASP) 7376 2.897 (ASP) 7376								
18487 26684 2.96 4.1E+00 UG7503.1 NT 17427 27641 2.31 4.1E+00 BE682425.1 EST_HUMAN 18138 2.89 4.1E+00 PO9716 SWISSPROT 18216 1.246 4.1E+00 BE885880.1 EST_HUMAN 19231 26285 1.37 4.0E+00 D47876 SWISSPROT 18648 28936 2.17 4.0E+00 P07564 SWISSPROT 18548 28936 2.47 3.9E+00 A64518.1 NT 14124 0.99 3.9E+00 A76546.1 NT 14324 2.47 3.9E+00 A76548.1 NT 15391 25454 2.47 3.9E+00 A7654 SWISSPROT 16391 25454 2.47 3.9E+00 A7654 SWISSPROT 16391 25454 2.47 3.9E+00 A7654 SWISSPROT 16391 25454 2.47 3.9E+00 A7650 SWISSPROT 16218	Probe SEQ ID NO:		ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
17427 Z7641 2.31 4.1E+00 BF692425.1 EST_HUMAN 18216 12.46 4.1E+00 P09716 SWISSPROT 18216 12.46 4.1E+00 BE885880.1 EST_HUMAN 1821 25241 1.86 4.1E+00 P47876 SWISSPROT 16131 26285 1.37 4.0E+00 P7876 SWISSPROT 16131 26285 1.37 4.0E+00 P7564 SWISSPROT 18348 28836 3.34 4.0E+00 P07564 SWISSPROT 18375 23181 3.8E+00 P664518.1 NT 14124 0.99 3.9E+00 P67564 SWISSPROT 15391 25456 2.47 3.9E+00 P6644357.1 EST_HUMAN 16200 28640 A.6E+00 P7564 SWISSPROT 16391 26443 3.9E+00 P6644357.1 EST_HUMAN 16391 28640 AR6644357.1 EST_HUMAN 16719 26812 3.9E+00	6817			2.95	4.1E+00		LV	Pan troglodytes novel repetitive solo LTR element in the RNU2 locus
18139 2.89 " 4.1E+00 P09716 SWISSPROT 18216 12.46 4.1E+00 BE8B5890.1 EST HUMAN 19231 25241 1.86 4.1E+00 P47878 SWISSPROT 16131 25285 1.37 4.0E+00 O33010 SWISSPROT 17917 28184 2.17 4.0E+00 P07564 SWISSPROT 18648 28934 3.34 4.0E+00 P07564 SWISSPROT 13375 23181 3.39 3.9E+00 P07564 SWISSPROT 15391 25456 2.47 3.9E+00 P07564 SWISSPROT 15391 25456 2.47 3.9E+00 P07564 SWISSPROT 16002 25456 2.47 3.9E+00 P07564 SWISSPROT 16012 25456 2.47 3.9E+00 P07564 SWISSPROT 16012 25456 2.47 3.9E+00 P07564 SWISSPROT 16012 25456 2.47 3.9E+00 P07564 SWISSP	7576		27641	2.31	4,1E+00		EST_HUMAN	602247938F1 NIH_MGC_62 Homo sapiens cDivA clone IMAGE:4333209 5'
18216 1246 4.1E+00 BE885890.1 EST_HUMAN 18231 25241 1.86 4.1E+00 P47876 SWISSPROT 16131 26285 1.37 4.0E+00 P7564 SWISSPROT 18648 28934 3.34 4.0E+00 P07564 SWISSPROT 18548 28934 3.34 4.0E+00 P07564 SWISSPROT 13375 23181 3.9E+00 X64318.1 NT 14124 25456 2.47 3.9E+00 R645686.1 NT 15391 25456 2.47 3.9E+00 R645686.1 NT 16002 26140 4.46 3.9E+00 R64318.1 NT 16218 26879 2.47 3.9E+00 R64366.1 NT 16719 26913 2.3E+00 R78696.1 NT 16719 26913 2.3E+00 R78696.1 NT 17879 28120 3.9E+00 R78696.1 NT 18680 28816 5.68	8228			2.89	4.1E+00		SWISSPROT	HYPOTHETICAL PROTEIN HVLF1
18648 28936 3.34 4.0E+00 PA7876 SWISSPROT 18648 28936 3.34 4.0E+00 P07564 SWISSPROT 1874 0.99 3.9E+00 X64518.1 NT 16002 25456 2.47 3.9E+00 AF055486.1 NT 16719 25646 2.47 3.9E+00 MF055486.1 NT 16719 25810 4.46 3.9E+00 MF055486.1 NT 16719 25810 4.46 3.9E+00 MF055486.1 NT 17879 28120 3.9E+00 MF05016.1 NT 17879 28120 3.9E+00 MF00165.1 NT 16789 28816 5.68 3.9E+00 MF00165.1 EST_HUMAN	8338			12.46	4.1E+00		EST_HUMAN	601507510F1 NIH_MGC_71 Hamo sapiens cDNA clone IMAGE:3909051 5'
16131 26285 1.37 4.0E+00 033010 SWISSPROT 17917 28184 2.17 4.0E+00 P07564 SWISSPROT 18648 28936 3.34 4.0E+00 P07564 SWISSPROT 18648 28936 3.34 4.0E+00 P07564 SWISSPROT 13375 23181 3.8E+00 AF055468.1 NT 14124 0.99 3.9E+00 AF055468.1 NT 15391 25454 2.47 3.9E+00 BE814357.1 EST_HUMAN 16702 253140 4.46 3.9E+00 P07564 SWISSPROT 16719 25455 2.47 3.9E+00 P07564 SWISSPROT 16710 25454 2.47 3.9E+00 P07564 SWISSPROT 16710 25455 2.47 3.9E+00 P07564 SWISSPROT 16710 25455 2.47 3.9E+00 P07564 SWISSPROT 16710 25913 3.9E+00 P07564 SWISSPROT	9672			1.86	4.1E+00		SWISSPROT	INSULIN-LIKE GROWTH FACTOR BINDING PROTEIN 1 PRECURSOR (IGFBP-1) (IBP-1) (IGF- BINDING PROTEIN 1)
17917 28164 2.17 4.0E+00 P14546 SWISSPROT 18648 28934 3.34 4.0E+00 P07564 SWISSPROT 18648 28936 3.34 4.0E+00 P07564 SWISSPROT 13375 223181 3.8E+00 AF055468.1 NT 14124 0.99 3.9E+00 AF055468.1 NT 15391 25454 2.47 3.9E+00 BE814357.1 EST_HUMAN 16391 25455 2.47 3.9E+00 BE814357.1 EST_HUMAN 16719 25913 2.3E+00 MZ3909.1 NT 16719 25913 3.9E+00 MZ3907.1 NT 16719 25913 3.9E+00 MZ3907.1 NT 16719 26913 3.9E+00 MZ3907.1 NT 16719 26913 3.9E+00 MZ3907.1 NT 16719 26120 3.9E+00 MZ3907.1 NT 1853 2681 3.9E+00 AF091439.1 EST_HUMAN </td <td>8266</td> <td></td> <td>26285</td> <td>1.37</td> <td>4.0E+00</td> <td></td> <td>SWISSPROT</td> <td>CELL DIVISION PROTEIN FTSY HOMOLOG</td>	8266		26285	1.37	4.0E+00		SWISSPROT	CELL DIVISION PROTEIN FTSY HOMOLOG
18648 Z8934 3.34 4.0E+00 P07564 SWISSPROT 18648 28935 3.34 4.0E+00 P07564 SWISSPROT 13375 223181 3.8E+00 X64518.1 NT 14124 0.39 3.9E+00 X64518.1 NT 15391 25454 2.47 3.9E+00 BE814357.1 EST HUMAN 15391 25455 2.47 3.9E+00 BE814357.1 EST HUMAN 16702 26913 2.47 3.9E+00 BE814357.1 EST HUMAN 16719 26913 2.32 3.9E+00 BE814357.1 INT 16719 26913 2.32 3.9E+00 BE814357.1 INT 16719 26913 2.32 3.9E+00 M23907.1 NT 16719 26913 2.32 3.9E+00 X65965.1 NT 16719 26913 2.4 3.9E+00 X66406.2 NT 16719 26923 3.9E+00 X66966.1 NT	8768	Ш		217	4.0E+00		SWISSPROT	CYTOCHROME C OXIDASE POLYPEPTIDE III
18648 228936 3.34 4.0E+00 P07564 SWISSPROT 13375 22181 3.89 3.9E+00 X64518.1 NT 14124 0.99 3.9E+00 X64518.1 NT 15391 25454 2.47 3.9E+00 BE814357.1 EST_HUMAN 15391 25456 2.47 3.9E+00 BE814357.1 EST_HUMAN 16002 28140 4.46 3.9E+00 P38299 SWISSPROT 16719 26379 4.66 3.9E+00 M23807.1 NT 16719 26312 2.32 3.9E+00 X65865.1 NT 17879 28120 3.9E+00 X78000.1 NT 17879 28120 3.9E+00 X78000.1 NT 18532 28816 5.68 3.9E+00 X661489.1 EST_HUMAN 18789 28822 1.18 3.7E+00 A461489.1 EST_HUMAN 18586 28872 2.13 3.7E+00 BF669279.1 EST_HUMAN	8835	_		3,34	4.0E+00		SWISSPROT	GENOME POLYPROTEIN [CONTAINS: CAPSID PROTEIN C (CORE PROTEIN); MATRIX PROTEIN (ENVELOPE GLYCOPROTEIN M); MAJOR ENVELOPE PROTEIN E; NONSTRUCTURAL PROTEINS NS1, NS2A, NS2B, NS4A AND NS4B; HELICASE (NS3); RNA-DIRECTED RNA POLYMERASE (NS5)]
18648 28936 3.34 4.0E+00 P07564 SWISSPROT 13375 23181 3.89 3.9E+00 X64518.1 NT 14124 25454 2.47 3.9E+00 R655468.1 NT 15391 25454 2.47 3.9E+00 BE814357.1 EST_HUMAN 15391 25455 2.47 3.9E+00 P38299 SWISSPROT 16002 253140 4.46 3.9E+00 P38299 SWISSPROT 16719 25913 2.32 3.9E+00 P38299 SWISSPROT 16719 25913 2.32 3.9E+00 P38299 SWISSPROT 16719 25913 2.32 3.9E+00 P38299 SWISSPROT 17879 28120 3.9E+00 P465885.1 NT 17879 28120 3.9E+00 P4661489.1 EST_HUMAN 18589 2892 1.18 3.8E+00 P4725.1 NT 18789 28872 1.18 3.7E+00 P4725.1 EST_HUMAN 18586 28872 2.13 3.7E+00 P4669279.1 EST_HUMAN 16899 27090<	_							GENOME POLYPROTEIN [CONTAINS: CAPSID PROTEIN C (CORE PROTEIN); MATRIX PROTEIN (ENVELOPE GLYCOPROTEIN M); MAJOR ENVELOPE PROTEIN E; NONSTRUCTURAL PROTEINS
14124 0.99 3.9E+00 AF055466.1 IT 15391 25454 2.47 3.9E+00 BE814357.1 EST_HUMAN 16002 26140 4.46 3.9E+00 BE814357.1 EST_HUMAN 16218 28379 4.68 3.9E+00 BS814357.1 EST_HUMAN 16719 28913 2.32 3.9E+00 M33907.1 NT 17879 28120 3.09 3.9E+00 M33907.1 NT 18532 2.8E+00 M33907.1 NT NT 18683 2.8E+00 M33907.1 NT NT 1878 2.8E+00 M33907.1 NT NT 1878 2.8E+00 M33907.1 NT NT 1878 2.8E+00 M33907.1 NT NT 18858 2.8E+00	3459	L_		3.89	3 9F+00		SWISSPROT	NS1, NS2A, NS2B, NS4A AND NS4B; HELICASE (NS3); RNA-DIRECTED RNA POLYMERASE (NS5)] N tabacum chilinase gene 50 for class I chilinase C
15391 25454 2.47 3.9E+00 BE814357.1 EST_HUMAN 15381 25455 2.47 3.9E+00 BE814357.1 EST_HUMAN 16002 28140 4.46 3.9E+00 MZ3807.1 NT 16719 28913 2.32 3.9E+00 MZ3807.1 NT 16719 28913 2.32 3.9E+00 MZ3807.1 NT 17879 28120 3.9E+00 MZ3807.1 NT 18532 2.8B16 5.69 3.9E+00 MZ3807.1 NT 12459 2.24 3.8E+00 MZ3807.1 NT 15789 2.8B16 5.69 3.9E+00 MZ3807.1 NT 16789 2.8B2 1.18 3.8E+00 MZ3807.1 NT 18586 25822 10.09 3.7E+00 MZ380.7 NT 18586 28872 2.13 3.7E+00 MZ392.9 NT 10516 2.23 3.6E+00 MZ380279.1 EST_HUMAN 18586 28872 2.13 3.7E+00 MZ392.9 RT_HUMAN 10516 27090 3.6E+00 MZ447.1 NT <td>4226</td> <td>_</td> <td></td> <td>0.99</td> <td>3.9E+00</td> <td>AF055468.1</td> <td>5</td> <td>Mus musculus seminal vesicle secretory protein 99 (MSVSP99) gene, promoter region</td>	4226	_		0.99	3.9E+00	AF055468.1	5	Mus musculus seminal vesicle secretory protein 99 (MSVSP99) gene, promoter region
15381 25455 2.47 3.9E+00 BE814357.1 EST_HUMAN 16002 28140 4.46 3.9E+00 P38299 SWISSPROT 16719 26913 2.32 3.9E+00 M23807.1 NT 17879 28120 3.09 3.9E+00 A661489.1 NT 18532 28816 5.69 3.9E+00 A661489.1 RT HUMAN 12459 2.4 3.8E+00 A661489.1 EST HUMAN 16789 2.6882 1.18 3.8E+00 A661489.1 EST HUMAN 18586 28872 1.18 3.8E+00 A661489.1 EST HUMAN 18586 28872 1.18 3.8E+00 A4725.1 EST HUMAN 18586 28872 2.13 3.7E+00 BF669279.1 EST HUMAN 10516 20322 2.13 3.7E+00 BF669279.1 EST HUMAN 10616 20322 2.13 3.6E+00 AV761056.1 EST HUMAN 10889 27090 3.66 3.6E+00 AV761056.1 EST HUMAN	6471	•		2.47	3.9E+00	BE814357.1	EST HUMAN	MR0-BN0070-300500-028-h05 BN0070 Homo saplens cDNA
16002 26140 4.46 3.9E+00 P38299 SWISSPROT 16218 26913 2.32 3.9E+00 M23807.1 NT 17879 26913 2.32 3.9E+00 A65865.1 NT 17879 28120 3.09 3.9E+00 Y18000.1 NT 12459 2.8 3.9E+00 A661489.1 EST_HUMAN 12459 2.4 3.8E+00 A661489.1 EST_HUMAN 16789 2.8982 1.18 3.8E+00 D44725.1 EST_HUMAN 18586 28872 2.13 3.7E+00 AL161539.2 NT 1656 2.2873 2.13 3.7E+00 BF669279.1 EST_HUMAN 1656 2.2873 2.13 3.7E+00 BF669279.1 EST_HUMAN 16516 2.0322 2.13 3.7E+00 BF669279.1 EST_HUMAN 16829 27090 3.6E+00 AV761055.1 EST_HUMAN 16829 27091 3.6E+00 AV761055.1 NT	5471	Ш	25455	2.47	3.9E+00		EST_HUMAN	MRO-BN0070-300500-028-h05 BN0070 Homo sapiens cDNA
16218 26379 4.68 3.8E+00 MZ3807.1 NT 16719 26913 2.32 3.8E+00 X65885.1 NT 17879 28120 3.09 3.8E+00 Y18000.1 NT 18532 28816 5.68 3.9E+00 A661489.1 EST_HUMAN 12459 2.4 3.8E+00 A661489.1 EST_HUMAN 16789 2882 1.18 3.8E+00 A661489.1 EST_HUMAN 18586 28872 1.18 3.8E+00 A660489.1 EST_HUMAN 18586 28872 2.13 3.7E+00 AL161539.2 NT 10516 20322 2.13 3.7E+00 BF669279.1 EST_HUMAN 10516 20322 2.13 3.7E+00 BF669279.1 EST_HUMAN 10516 20322 2.1 3.6E+00 AV761056.1 EST_HUMAN 16899 27091 3.6E+00 AV761056.1 NT	6108			4.46	3.9E+00		SWISSPROT	HYPOTHETICAL TRANSCRIPTIONAL REGULATOR IN AIDB-RPSF INTERGENIC REGION
16719 26913 2.32 3.8E+00 (X65895.1 NT 17879 28120 3.09 3.8E+00 (Y18000.1 NT 18532 28816 5.68 3.9E+00 (Y18000.1 NT 12459 2.4 3.8E+00 (A661489.1) EST HUMAN 16789 28822 1.18 3.8E+00 (A4725.1) EST HUMAN 18586 28872 2.13 3.7E+00 (A161639.2) NT 1658 28873 2.13 3.7E+00 (A761055.1) EST HUMAN 16899 27090 3.66 3.6E+00 (A761055.1) EST HUMAN 16899 27091 3.66 3.6E+00 (A761055.1) EST HUMAN 16899 27091 3.66 3.6E+00 (A761055.1) NT	6355	l		4.68	3.9€+00		<u>1</u> 21	Human MHC class II lymphocyte antigan (DPw4-beta-1) gene, exon 2
17879 28120 3.09 3.8E+00 Y18000.1 NT 18532 28816 5.68 3.9E+00 A661489.1 EST_HUMAN 12459 2.4 3.8E+00 A661489.1 EST_HUMAN 16789 26922 1.18 3.8E+00 D4725.1 EST_HUMAN 18586 28872 2.13 3.7E+00 BF669279.1 EST_HUMAN 16516 20322 2.13 3.7E+00 BF669279.1 EST_HUMAN 16899 27090 3.66 3.6E+00 AV761055.1 EST_HUMAN 16899 27091 3.66 3.6E+00 AV761055.1 NT 16899 27091 3.66 3.6E+00 AV761055.1 NT	88 849			2:32	3.9E+00		Ę	X.laevis mRNA for M4 muscarinic receptor
18532 28816 5.68 3.9E+00 AA661489.1 EST_HUMAN 12459 2.4 3.8E+00 AE001652.1 NT 16789 2.6882 1.18 3.8E+00 D44725.1 EST_HUMAN 13846 25622 10.09 3.7E+00 AL161539.2 NT 18586 22872 2.13 3.7E+00 BF669279.1 EST_HUMAN 10516 20322 2.1 3.6E+00 AV761055.1 EST_HUMAN 16899 27090 3.66 3.6E+00 AV761055.1 EST_HUMAN 16899 27091 3.6E+00 AV761055.1 NT	8695			3.09	3.9E+00		F	Homo saplens NF2 gene
12459 2.4 3.8E+00 AE001662.1 NT 16789 25982 1.18 3.8E+00 D44725.1 EST_HUMAN 13846 23622 10.09 3.7E+00 AL161539.2 NT 18586 28872 2.13 3.7E+00 BF669279.1 EST_HUMAN 10516 20322 2.1 3.7E+00 BF669279.1 EST_HUMAN 16899 27090 3.6E 3.6E+00 AV761055.1 EST_HUMAN 16899 27091 3.6E 3.6E+00 AV761055.1 NT 16899 27091 3.6E 3.6E+00 AV761055.1 NT	8715			5.68	3.9E+00		EST_HUMAN	nr18a12.s1 NCI_CGAP_Ew1 Homo saplens cDNA clone IMAGE:1168318 similar to gb:A10416 METALLOPROTEINASE INHIBITOR 1 PRECURSOR (HUMAN);
16789 26982 1.18 3.8E+00 D44725.1 EST_HUMAN 13846 23622 10.09 3.7E+00 AL161639.2 NT 18586 28872 2.13 3.7E+00 BF669279.1 EST_HUMAN 18586 28873 2.13 3.7E+00 BF669279.1 EST_HUMAN 10516 20322 2.1 3.6E+00 AV761055.1 EST_HUMAN 16899 27090 3.66 3.6E+00 AV761055.1 NT 16899 27091 3.66 3.6E+00 AV761057.1 NT	2588			2.4	3.8E+00		Þ	Helicobacter pylori, strain J99 section 123 of 132 of the complete genome
13846 23622 10.09 3.7E+00 AL161639.2 NT 18586 28872 2.13 3.7E+00 BF669279.1 EST_HUMAN 10516 20322 2.13 3.7E+00 BF669279.1 EST_HUMAN 10616 20322 2.1 3.6E+00 AV761055.1 EST_HUMAN 16899 27090 3.6B 3.6E+00 AV761055.1 NT 16899 27091 3.6B 3.6E+00 AV761057.1 NT	6911			1.18	3.8E+00		EST_HUMAN	HUMSUPY135 Human brain cDNA Homo saplens cDNA clone 148
18586 28872 2.13 3.7E+00 BF669279.1 EST_HUMAN 18586 28873 2.13 3.7E+00 BF669279.1 EST_HUMAN 10616 20322 2.1 3.6E+00 AV761055.1 EST_HUMAN 16899 27090 3.66 3.6E+00 AE00447.1 NT 16899 27091 3.66 3.6E+00 AE00447.1 NT	3936			10.09	3.7E+00		Z-L	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 39
18586 28873 2.13 3.7E+00 BF669279.1 EST_HUMAN 10616 20322 2.1 3.6E+00 AV761055.1 EST_HUMAN 16899 27090 3.66 3.6E+00 AE00447.1 NT 16899 27091 3.66 3.6E+00 AE00447.1 NT	8730			2.13	3.7E+00		EST_HUMAN	602120551F1 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:4277748 5
10616 20322 2.1 3.6E+00 AV761055.1 EST_HUMAN 16899 27090 3.66 3.6E+00 AE00447.1 NT 16899 27091 3.66 3.6E+00 AE00447.1 NT	8730			2.13	3.7E+00	BF669279.1	EST_HUMAN	602120551F1 NIH_MGC_56 Homo sepiens cDNA clone IMAGE:4277748 5
16899 27090 3.66 3.6E+00 AE00447.1 NT 16899 27091 3.66 3.6E+00 AE00447.1 NT	678			2.1		AV761055.1	EST_HUMAN	AV761055 MDS Homo sapiens cDNA clone MDSBUE10 5'
16899 27091 3.66 3.6E+00 AE004447.1 (NT	7022			3.68			L/	Pseudomonas aeruginosa PA01, section 8 of 529 of the complete genome
	7022	╝		3.66	3.6E+00		卢	Pseudomonas æruginosa PA01, section 8 of 529 of the complete genome

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
8228	18110		3.78	3.6E+00	M96795.1	· FN	Escherichia coli glycerophosphata dehydrogenaise (glpD) gene, partial cds; and the translation start site has been verified (glpE), the translation start site has been verified (glpG), and repressor protein (glpR) genes, complete cds
3209	13133	22834	1.04	3.5E+00	AF221538.1	Ę	Cryptosporidium felis heat shock protein 70 (HSP70) gene, partial cds
1496	11400	21260	3.57	3.4E+00	AF254577.1	Þ	Brassica napus RPB5d mRNA, complete cds
2532	12408	22298	1.02	3.4E+00	AL163278.2	Ę	Homo sapiens chromosome 21 segment HS21¢078
6354	16217	26378	2.41		P04052	SWISSPROT	DNA-DIRECTED RNA POLYMERASE II LARGEST SUBUNIT
7889			3.17		AF013167.1	LN T	Saccharomyces cerevisiae MSS1 gene, complete cds
8818	18631	28919	1.92		L77570.1	NT	Homo saplens DiGeorge syndrome critical region, centromeric end
5036	14808	24679	1.41	3.3E+00	T662155 NT	Ŋ	Homo sapiens KIAA0480 gene product (KIAA0480), mRNA
5036	14908	24680	1.41	3.3E+00	7662155 NT	۲	Homo sapiens KIAA0480 gene product (KIAA0480), mRNA
492	10435	20248	1.39	3.2E+00	X96422.1	Į.	D.rerio zp-50 POU gene
3938	10435	20248	0.79	· 3.2E+00	X96422.1	L	D.reno 20-50 POU gena
5439	15359		2.68	3.2E+00	P12783	SWISSPROT	PHOSPHOGLYCERATE KINASE, CYTOSOLIC
5438	15359	25416	2.68		P12783	SWISSPROT	PHOSPHOGLYCERATE KINASE, CYTOSOLIC
9089	15713	25825	2.08	3.25+00	P18931	SWISSPROT	NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 4
5808	15713	22826	2.06	3.2E+00	P18931	SWISSPROT	NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 4
6269			2.35		Y13655.1	NT	Chlamydomonas reinhardtii chloroplast DNA for rps9, ycf4, ycf3, rps18 genes
6269		26610	2.35			NT	Chlamydomonas reinhardtii chloroplast DNA for rps9, ycf4, ycf3, rps18 genes
7221	17098		6.33	3.2E+00	P13061	SWISSPROT	PERIPLASMIC [NIFE] HYDROGENASE SMALL SUBUNIT (NIFE HYDROGENLYASE SMALL CHAIN)
7491	17381	27567	1.17	3.2E+00	M36383.1	NT	S.cerevisiae threonine deaminase (ILV1) gene, complete cds
. 7837	17687	22822	1.69	3.2E+00	AB016081.2	NT	Oryzlas latipes OIGC6 gene for guanylyl cyclase C, complete cds
808	18863		4.32	3.2E+00	1.33836.1	TN	Sus scrofa choline acetyltransferase gene, promoter region
285	15497	25574	2.42	3.1E+00	Q10135	SWISSPROT	HYPOTHETICAL 142.5 KD PROTEIN C23E2.02 IN CHROMOSOME I
7000	16877	27067	4.35	3.1E+00	P49894	SWISSPROT	TYPE I IODOTHYRONINE DEIODINASE (TYPE-I 5'DEIODINASE) (DIOI) (TYPE 1 DI) (5DI)
7000	16877	89022	4.35	3.1E+00	P49894	SWISSPROT	TYPE I IODOTHYRONINE DEIODINASE (TYPE-I 5'DEIODINASE) (DIOI) (TYPE 1 DI) (5DI)
7334	17238		3.8	3.1E+00	Q14957	SWISSPROT	GLUTAMATE [NMDA] RECEPTOR SUBUNIT EPSILON 3 PRECURSOR (N-METHYL D-ASPARTATE RECEPTOR SUBTYPE 2C) (NR2C) (NMDAR2C)
7920	17770	28009	4.78	3.1E+00	P49365	SWISSPROT	DEOXYHYPUSINE SYNTHASE (DHS)
							GENOME POLYPROTEIN (CONTAINS: CAPSID PROTEIN C (CORE PROTEIN); MATRIX PROTEIN CENVELORE DE DECTEIN DE PROTEIN DE L'ACHIET DAT DE LE NOMETEIN DE L'ACHIET DE L'ACHI
8759	17908		3.78	3.1E+00	P33515	SWISSPROT	NS2A, NS2B, NS4A AND NS4B, HELICASE (NS3), RNA-DIRECTED RNA POLYMERASE (NS5))
07774	70807		20.00	2 46	A Capan	TIV	retinoic acid nuclear receptor isoform beta 2 [mica, embryonal carcinoma cell line, PCC7-MZ1, mRNA, 2971
4//0	-1		0.30	3.15.400	330000.1) ii

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Table 4
Single Exon Probes Expressed in Heart

					<u> </u>		פווימים בעבודו ויסים בעלי מסים ווייסים וויסים וויס
Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acessian No.	Top Hit Database Source	Тф Hit Descriptor
2807	12737		1.06	3.0E+00	8923984 NT		Homo saplens hypothetical protein PRO0889 (FRO0889), mRNA
6273	15195	24970	1.83	3.0E+00	X53096.1	NT	S.aureus genes encoding Sau96i DNA methyltransferase and Sau96i restriction endonuclease
6245	ļ	1	9.6	3.0E+00	P18406	SWISSPROT	CYR61 PROTEIN PRECURSOR (3CH61)
7164	17041		1.45	3.0E+00	X67838.1	NT TN	B.napus DNA for myrosinase
8377	18251	20 R		4 DE	C7815d	TOROSSIME	RETINAL GUANYLYL CYCLASE 2 PRECURS OR (GUANYLATE CYCLASE 2F, RETINAL) (RETGC-2) (ROD OUTER SEGMENT MEMBRANE GUANYLATE CYCLASE 2) (ROS-GC2) (GUANYLATE CYCLASE P) (GC-F)
3	1					T	TO TOTAL TO THE TAX TO THE PROPERTY OF THE PRO
							RETINAL GUANYLYL CYCLASE 2 PRECURSOR (GUANYLATE CYCLASE 2F, KE I INAL) (KE I GC-2) (ROD OUTER SEGMENT MEMBRANE GUANYLATE CYCLASE 2) (ROS-GC2) (GUANYLATE CYCLASE
8374	18251	28502	6.51	3.0E+00	P51842	SWISSPROT	F) (GC.F)
1984	11858		2.58		AE002225.2	NT	Chlamydophila pneumoniae AR39, section 53 of 94 of the complete genome
6128	15976	26112	1.63	2.9E+00	236879.1	NT	F.pringlei gdcsPA gene for P-protein of the glycine cleavage system
6282	16146	26300	4.47	2.9E+00		SWISSPROT	BRAIN-SPECIFIC ANGIOGENESIS INHIBITOR 1 PRECURSOR
6282	16148	26301	4.47	2.9E+00			BRAIN-SPECIFIC ANGIOGENESIS INHIBITOR 1 PRECURSOR
6410	16271	26433	5.19	2.9€+00		ISSPROT	ADHERENCE FACTOR (ADHESION AND AGGREGATION MEDIATING SURFACE ANTIGEN)
1441	11346	21212	6.79	2.8E+00	AF186398.1	NT	Buxus harlandii maturase K (matk) gene, partial cds; chloropiast gene for chloropiast product
1615	11519		3.12	2.8E+00	AL161552.2	본	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 52
6325					8393724 NT	TN	Mus musculus endomucin (LOC53423), mRNA
230	10199	20012	4.63	2.7E+00	TN 8088308 NT	NT	Mus musculus per-hexamer repeat gene 3 (Phx/3), mRNA
230	10199	20013	4.63	2.7E+00	6679306	L	Mus musculus per-hexamer repeat gene 3 (Phx;3), mRNA
5408	15327	25377	1.75	2.7E+00	L14005.1	NT	Homo sapiens apoA polymorphism Kringle IV gene, exons 1 and 2
7185	17062		2.21	2.7E+00	AL116459.1	NT	Botrytts cinerea strain T4 cDNA library under conditions of nitrogen deprivation
7996	17846		2.18	2.7E+00	BE063527.1	EST_HUMAN	CM0-BT0281-031199-037-h04 BT0281 Homo saplens cDNA
4578	14487	24253	4.35	2.6E+00	AF068749.1	Ŋ	Mus musculus sphingosine kinase (SPHK1b) mRNA, complete cds
5405	15324	25373		2.6E+00	IN 1095529	NT	Mus musculus SRY-box containing gene 13 (Sox13), mRNA
5405	15324	25374		2.6E+00	6755601 NT	ΙN	Mus musculus SRY-box containing gene 13 (Sox13), mRNA
6533	16391		5.42	2.6E+00	AF235502.1	NT	Mus musculus SH2-containing Inositol 5-phosphatase (Ship) gene, exons 16 through 27, and complete cds
9699	16576	26767	1.2	2.6E+00	AJ132180.1	NT	faba bean necrotic yellows virus C2-Eg gene, Isolate Egyptian EV1-93
9699	16576		1.2	2.6E+00	AJ132180.1	NT	faba bean necrotic yellows virus C2-Eg gene, Isolate Egyptian EV1-93
7567	17418	27634	2.85	. 2.6E+00	AL161540.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 40
7927	17777		1.52		8055193 NT	F	Mus musculus cleavage and polyadenylation specificity factor 3 (Cps/3), mRNA
9711					11419220	Ę	Homo sapiens ATP-binding cassette, sub-family B (MDR/TAP), member 4 (ABCB4), mRNA
1448	11353	21216	2.08	2.5E+00	AJ271844.1	占	Aspergillus nidulans recQ gene for DNA helicase, exons 1-4

PCT/US01/00666

WO 01/57274

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Table 4
Single Exon Probes Expressed in Heart

Single Exoll Flobes Expressed in near	Tcp Hit Descriptor	Aspergillus nidulans recQ gene for DNA helicase, exons 1-4	TEICHOIC ACID BIOSYNTHESIS PROTEIN F	QV4-FT0005-110500-205-g07 FT0005 Homo saplens cDNA	Rice DNA for aidolase C-1, complate cds	Mus musculus EIF4H gene, partial cds; LIMK1 gene, complete cds; and ELN gene, partial cds	Chicken alpha-3 collagen type VI mRNA, 3' end	Homo sapiens double C2-like domains, alpha (COC2A) mRNA	VITELLOGENIN I PRECURSOR (YOLK PROTEIN 1)	CD27L RECEPTOR PRECURSOR (T-CELL ACTIVATION ANTIGEN CD27) (T14)	CD27L RECEPTOR PRECURSOR (T-CELL ACTIVATION ANTIGEN CD27) (T14)	Helicobacter pylori, strain J99 section 47 of 132 of the complete genome	RC2-PT0004-031299-011-d05 PT0004 Homo sapiens cDNA	ENDOCHITINASE B PRECURSOR (CHN-B)	SKIN GRANULE PROTEIN PRECURSOR	SKIN GRANULE PROTEIN PRECURSOR	H:saplens CTGF gene and promoter region	XYLULOSE KINASE (XYLULOKINASE)	hre3f08.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:3133187 3'	hre3f06.x1 NCI_CGAP_Kld11 Homo sapiens cDNA clone IMAGE:3133187 3'	Bacillus subtilis chromosomal DNA, region 75 dégrees: glpPFKD operon and downstream	Fragaria x ananassa cytosolic ascorbate peroxidase (ApxSC) gene, ApxSC-c allele, complete cds	G.domesticus artificial single chain antibody gene (L3)	Bos taurus partial cytb gene for cytochrome b	Rattus norvegicus ATPase, Ca++ transporting, ibiquitous (Atp2a3), mRNA	MAJOR CENTROMERE AUTOANTIGEN B (CENTROMERE PROTEIN B) (CENP-B)	M.mazel dnaK and dnaJ genes homologues coding for DnaK and DnaJ	ALPHA(1,3)-FUCOSYLTRANSFERASE (GALACTOSIDE 3-L-FUCOSYLTRANSFERASE) (FUCOSYLTRANSFERASE 4) (FUCT-IV)	602069121F1 NIH_MGC_58 Homo saplens cDNA clone IMAGE:4068173 5'	602069121F1 NIH_MGC_58 Homo saplens cDNA clone IMAGE:4068173 5'			
וום בעחוו גוחחם	Top Hit Datebase Source	Г	SWISSPROT	SWISSPROT T	SWISSPROT	SWISSPROT T	EST_HUMAN C					SWISSPROT (V	SWISSPROT	SWISSPROT	NT			SWISSPROT S	SWISSPROT	H LN	SWISSPROT		EST_HUMAN h	B	- E	LN TN	NT PN		SWISSPROT	NT	A SWISSPROT	Г	П
	Top Hit Acession No.	14.1	P13485	P13485	P13485	P13485	AW949158.1	DE0307.1	5.1	M24282.1	4503352 NT	P02843	P26842	P26842	AE001486.1	AW875126.1	P24091	P13673	P13673	X92511.1	P09099	BE326702.1	BE326702.1	Y14079.1	AF158652.2	Z46724.1	.1	6978554 NT	P07199	X60265.1	011127	BF541987.1	BF541987.1
	Most Similar (Top) Hit BLAST E Value	2.5E+00	2.5E+00	2.5E+00	2.6E+00	2.5E+00	2.5E+00	2.6E+00		2.4E+00	2.4E+00	2.4E+00	2.4E+00	2.4E+00	2.4E+00			2.4E+00	2.4E+00		2.4E+00		2.4E+00	2.4E+00	2.4E+00	2.3E+00	2.3E+00		2.3E+00	2.3E+00	2.35+00		
	Expression Signal	2.08	2.33	2.33	1.71	1.71	1.34	1.76	2.28	0.86	7.62	4.19	2.14	2.14	2.63	1.71	9.45	25	2.5	2.18	6.49	1.67	1.67	1.77	2.52	9.36	1.28	2.22	2.3	1.53	181	2.6	2.6
	ORF SEQ ID NO:		25537			25538		27334				25664	26802	26803				£3872	27854	27911		28002	28003	28558	28839	20992		26431		28613	27364		Н
	Exan SEQ ID NO:	11353	15468	15468	15468	15468	16419	17141		12907	14699	15569	16613	16613	16659	16903	17002	17621	17621	17870	17731	17764	17764	18302	18554	11140	13934	16269	19766	16431	17165	1	1 1
	Probe SEQ ID NO:	1448	5552	6552	5886	2886	6561	7264	9086	2880	4816	5857	6733	6733	6780	7026	7125	7771	1777	7820	7881	7914	7914	8428	8665	1233	4031	6408	6479	6573	7289	8888	8888

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Top Hit Descriptor	601433673F1 NIH_MGC_72 Homo sepiens cDNA clone IMAGE:3918843 5	Neurospora crassa G protein alpha subunit GNA-3 (gna-3) gene, complete cds	Rat gene for regucalcin, exan1 (non-coding exon)	Rat gene for regucalcin, exon1 (non-coding exon)	SORTILIN-RELATED RECEPTOR PRECURSOR (SORTING PROTEIN-RELATED RECEPTOR CONTAINING LDLR CLASS A REPEATS) (MSORLA) (SORLA-1) (LOW-DENSITY LIPOPROTEIN RECEPTOR RELATIVE WITH 11 LIGAND-BINDING REPEATS) (LDLR RELATIVE WITH 11 LIGAND-BINDING REPEATS) (LAL1) (>	SORTILIN-RELATED RECEPTOR PRECURSOR (SORTING PROTEIN-RELATED RECEPTOR CONTAINING LDLR CLASS A REPEATS) (MGORLA) (SORLA-1) (LOW-DENSITY LIPOPROTEIN RECEPTOR RELATIVE WITH 11 LIGAND-BINDING REPEATS) (LDLR RELATIVE WITH 11 LIGAND-BINDING REPEATS)	BINDING REPEATS) (LR11) (>	60094340171 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:2959777 3'	MINOR VIRION STRUCTURAL PROTEIN MU:2	INSULIN-LIKE GROW TH FACTOR II PRECURSOR (IGF-II) (SOMATOMEDIN A)	nl95b02.s1 NCI_CGAP_Co10 Homo sapiens cDNA clone IMAGE:1058379 3'	zx05g10.r1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA clone IMAGE:785634 5'	601594733F1 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:3948561 5'	TRANSPOSON TY1 PROTEIN A	qm69b03.x1 Soares_placenta_8to9weeks_2NbHP8to9W Homo sapiens cDNA clone IMAGE:1893965 3' similar to gb:Y00433 GLUTATHIONE PEROXIDASE (HUMAN);	qm69b03.x1 Soares_placenta_8to8weeks_2NbHP8to9W Homo saplens cDNA clone IMAGE:1893965 3' similar to gb:Y00433 GLUTATHIONE PEROXIDASE (HUMAN);	601855591F1 NIH_MGC_57 Homo sapiens cDNA done IMAGE:4076391 6'	Homo sapiens ovarian granulosa cell 13.0 kDa protein hGR74 homotog mRNA, complete cds	UROMODULIN PRECURSOR (TAMM-HORSFALL URINARY GLYCOPROTEIN) (THP)	EARLY E1A 28 KD PROTEIN	Mus musculus pre-T cell receptor alpha gene, eithancer region and upstream region	UFH-BI3-aki-e-08-0-UI.s1 NCI_CGAP_Sub5 Homo sapiens cDNA clone IMAGE:2734550 3'	ALPHA-2-HS-GLYCOPROTEIN PRECURSOR (FETUIN-A)	yy08a10.s1 Soares melanocyte 2NbHM Horno signens cDNA clone IMAGE:270818 3' similar to gb:M55854 TRANSCRIPTION INITIATION FACTOR TFIID (HUMAN);	AU123630 NT2RM2 Homo sapiens cDNA clone NT2RM2000671 5'	Dome continue applicable (PDOVDEL) mONA commission and
Top Hit Database Source	EST_HUMAN 6	N N	NT.	NT	SWISSPROT B	0 PE	SWISSPROT & B	EST_HUMAN 6	SWISSPROT IN	SWISSPROT	EST_HUMAN n	EST_HUMAN z	EST_HUMAN 6	SWISSPROT T	EST_HUMAN s	EST HUMAN S	EST_HUMAN 6	F	SWISSPROT	ISSPROT		EST_HUMAN U	SWISSPROT	EST HUMAN T	Г	T
Top Hit Acession No.	BE895237.1	AF281862.1	D67071.1	D67071.1	208880		088307	BE250383.1	000335	P51459	AA594574.1	AA449012.1	BE741678.1	Q04706	AI290373.1	AI290373.1	BF246782.1	AF183416.1	P07911	P10407	AF132612.2	AW449366.1	070159	N29575.1	AU123630.1	A E400827 4
Most Similar (Top) Hit BLAST E Value		2.3E+00	2.2E+00		2.2E+00			2.2E+00	2.2E+00			2.2E+00	2.2E+00			2.2E+00	2.2E+00	2.2E+00	2.2E+00	2.2E+00	2.1E+00	2.1E+00	2.1E+00	2.1E+00	2.1E+00	
Expression Signal	4	1.37	3.82	3.82	10.08		10.08	90.6	3.3	2.89	3.14	51.56	11.83	2.1	1.56	1.56	2.28	2.88	4.5	4.67	6.81	1.19	3.72	4.7	1.89	4 23
ORF SEQ ID NO:	25333		23891		24873		24974		25853	26005		26514			27865	27866		28066	28132	56682	20304		26208	26087		2002
Exan SEQ ID NO:	19002	19405	14114	14114	15198		15198	15605	15741	15882	15113		17250	19468	17633	17633	17662	17825	17888	18700		13465	16057	15955	1	ı
Probe SEQ ID NO:	9304	8838	4216	4216	5276		5276	9699	5835	5978	6155	6486	7381	7518	7783	7783	7812	7975	8739	8889	557	3539	6074	6195	6948	1178

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Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
9428	19584		4.01	1.8E+00	AF314254.1	FA	Chlamydomonas reinhardtii alternative oxidase 1 (AOX1) gene, nuclear gene encoding mitochondrial protein
9504	19119		2.4	1.8E+00	9506404 NT	Ę	Rattus norvegicus Actin-related protein complex (1b (Arpc1b), mRNA
8988	19429		1.34	1.8E+00	BF316805.1	EST_HUMAN	601903309F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE 4135586 5'
1092	11008		1.92	1.7E+00	Q60114	SWISSPROT	LEVANSUCRASE (BETA-D-FRUCTOFURANOSYL TRANSFERASE) (SUCROSE 6-FRUCTOSYL TRANSFERASE)
2225	12110	22013	3.25		AL163280.2	닏	Homo sapiens chromosome 21 segment HS21C080
2321	12202		1.02	1.7E+00	AI141067.1	EST_HUMAN	oz43h05.x1 Soares_NhHMPu_S1 Homo sapiens cDNA clone IMAGE:1678137 3'
4356	14252	24037	0.84	1.7E+00	•	SWISSPROT	LEVANSUCRASE (BETA-D-FRUCTOFURANOSYL TRANSFERASE) (SUCROSE 6-FRUCTOSYL TRANSFERASE)
6448	15369	25424	1.48	1.7E+00	BE063548.1	EST_HUMAN	CM0-BT0282-171299-127-e05 BT0282 Homo sapiens cDNA
6448	15369	25425	1.48	1.7E+00	BE063546.1	EST_HUMAN	CM/0-BT0282-171299-127-e05 BT0282 Homo saplens cDNA
5664			3.58	1.7E+00	Q9TTR8	SWISSPROT	COUP TRANSCRIPTION FACTOR 1 (COUP-7F1) (COUP-TF1)
7014			1.29	1.7E+00	BF308000.1	EST_HUMAN	601894255F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:4140084 5
7282	19467	27355	2.15	1.7E+00	060479	SWISSPROT	HOMEOBOX PROTEIN DLX3
7282	19467	27356	2.15	1.7E+00	١	SWISSPROT	HOMEOBOX PROTEIN DLX-3
8874	18686	28977	1.78	1.7E+00	W22424.1	EST_HUMAN	67B7 Human retina cDNA Tsp5091-cleaved sublibrary Homo saplens cDNA not directional
8384	19044	25306	137	1 7F+00	A1678443 1	EST HIMAN	tu82d07.x1 NCI_CGAP_Gas4 Homo saplens cDNA clone IMAGE:2257549 3' similar to contains MSR1.t1 MSR1 repetitive element:
	L						qf50b01.x1 Soares testis NHT Homo sapiens cDNA clone IMAGE:1753417 3' similar to contains L1.t1 L1
9860	19351	25185	2.26	1.7E+00	Al198573.1	EST_HUMAN	repatitive element;
1989	11882	21775	16.73	1.6E+00	AF199339.1	NT	Homo sapiens lens epithelium-derived growth factor gene, alternatively spliced, complete cds
1897	11891	24783	3.61	1.6E+00	AF077374.1	NT	Homo sapiens small proline-rich protein (SPRR3) gene, exons 1, 2, and 3 and complete cds
2003	11896	21788	1.96	1.6E+00	Y11344.1	NT	Mus musculus ST6GalNAcill gene, exon 2
2238	12122		1.48	1.6E+00	X98373.1	NT	B.napus gene encoding endo-polygalacturonase
2034	42868	22858	080	4 65 400	W/59428 1	DOT LIMAN	zd25f01.r1 Soares_fetal_heart_NbHH19W Homb septens cDNA clone IMAGE:341689 5' similar to
3946	L		5.14		BF570077.1	EST HUMAN	602186095T1 NIH MGC 45 Homo saplens cDNA clone IMAGE:4310591 3
4255	1	23928	1.44		AF155827.1	E	Homo saplens proliferation-associated SNF2-like protein (SMARCA6) mRNA, complete cds
4265	14154	23929	1.44	1.6E+00	AF155827.1	Ę	Homo sapiens proliferation-associated SNF2-like protein (SMARCA6) mRNA, complete cds
5016	14890	24657	3.14	1.6E+00	Y11344.1	NT	Mus musculus ST6GalNAclil gene, exon 2
5016				1.6E+00	Y11344.1	NT	Mus musculus ST6GalNAcill gene, exon 2
5558	_		2.19	1.6E+00	L04808.1	i	Brachydanio reno MHC class II DA-beta-2*01 gene, 3' end
6296	16160	28317	2.84	1.6E+00	BE697267.1	EST_HUMAN	RC0-CT0415-200700-032-c10 CT0415 Homo sapiens cDNA

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Probe Exon SEQ ID SEQ ID NO: NO:	ORF SEQ ID ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acesslon No.	Top Hit Database Source	Ττφ Hit Descriptor
6681 16	16561	1.21	1.6E+00	Q46378	SWISSPROT	VIRULENCE FACTOR MVIN HOMOLOG
6881 16	16760 26958	3.56	1.6E+00	AJ297131.1	NT	Mus musculus SIL, MAP_17, CYP_a, SCL & CYP_b genes
7444 19	19465 26646	1.3	1.6E+00	X52046.1	NT	M.musculus COL3A1 gene for collegen alpha-1
7444 19	19465 26647	1.3	1.6E+00		NT	M.musculus COL3A1 gene for collegen alpha-1
7611 17	17462 27679	1.29	1.6E+00		EST_HUMAN	ph6b6_19/1TV Outward Alu-primed hncDNA library Homo sapiens cDNA clone ph6b6_19/1TV
	17702 27946	1.25	1.6E+00		EST_HUMAN	QV4-LT0016-090200-100-d07 LT0016 Homo saplens cDNA
7862 17	17702 27947	1.26	1.6E+00	AW835644.1	EST_HUMAN	QV4-LT0016-090200-100-d07 LT0016 Homo sapiens cDNA
8217 15	15520 25601	5.86	1.6E+00	AF005631.1	NT	Homo sapiens transglutaminase type I (Tgasel) gene, promoter region
8944 18	18752 29047	3.25	1.6E+00	AF104313.1	NT	Нотто sapiens unknown mRNA
9902 193	18379	1.65	1.6E+00	AV764043.1	EST_HUMAN	AV764043 MDS Homo saplens cDNA clone MDSDAH08 5'
30 100	10017 19812	4.29	1.5E+00	U53449.1	NT	Rettus norvegicus jun dimerization protein 2 (jdp-2) mRNA, complete cds
231 103	10200 20014	1.76	1.5E+00	AE002201.2	NT	Chlamydophila pneumoniae AR39, section 32 of 94 of the complete genome
605 109	10541	1.79	1.5E+00	6752961 NT	5	Mus musculus a disintegrin and metalloproteinase domain (ADAM) 15 (metargidin) (Adam15), mRNA
2359 12	12239 22135	2.46	1.5E+00	AJ131402.1	NT	Potato virus A RNA complete genome, Isolate U
2468 12	12344 22237	2.02	1.5E+00	6678350 NT	NT	Mus musculus T-cell lymphoma invasion and matastasis 1 (Tiam1), mRNA
3099 12	12239 22135	2.22	1.5E+00	AJ131402.1	NT	Potato virus A RNA complete genome, isolate U,
3329 13;	13249 23054	0.82	1.5E+00	AE001945.1	NT	Deinococcus radiodurans R1 section 82 of 229 of the complete chromosome 1
5856 15	15762 25880	2.71	1.5E+00	R17879.1	EST_HUMAN	yg10e02.r1 Soares infant brain 1NIB Homo sapiens cDNA clone IMAGE:31693 5'
6230 16	16096	1.42	1.5E+00	BE785356.1	EST_HUMAN	601478745F1 NIH_MGC_68 Homo sapiens cDINA clone IMAGE:3881655 5'
			1.5E+00		SWISSPROT	HYPOTHETICAL 118.4 KD PROTEIN IN BAT2-DAL5 INTERGENIC REGION PRECURSOR
6250 16	16116 26269	29.13	1.5E+00	P47179	SWISSPROT	HYPOTHETICAL 118.4 KD PROTEIN IN BATZ-DALS INTERGENIC REGION PRECURSOR
7671 178	17521 27747	7.58	1.5E+00		EST HUMAN	RC0-TN0078-150900-034-g05 TN0078 Homo sapiens cDNA
7782 170	17632	1.7.1	1.5E+00	BF337944.1	EST_HUMAN	602035771F1 NCL_CGAP_Bm64 Homo sapiens cDNA clone IMAGE:4183865 5'
7856 17	17706 27951	1.96	1.5E+00		EST_HUMAN	ze38g06.r1 Soares retina N2b4HR Homo sapiens cDNA clone IMAGE:361306 5'
7856 17	17706 27952	1.96	1.5E+00		EST_HUMAN	ze38g06.r1 Soares retina N2b4HR Homo sapiens cDNA clone IMAGE:381306 5'
8702 18	18520 28802		1.5E+00	AL134197.1	EST_HUMAN	DKFZp547P243_s1 547 (synonym: hfbr1) Homo sapiens cDNA clone DKFZp547P243 3'
8828 18	18641	10.73	1.5E+00	X07380.1	NT	Matze mitochondrial tRNA-Ser gene and tRNA-Phe pseudogene
9223 18(18946	1.6		6763287 NT	NT	Mus musculus caspase 8 associated protein 2 (Casp8ap2), mRNA
9596 19	19178	2.89	1.5E+00	AL445065.1	NT	Thermoplasma acidophilum complete genome; segment 3/5
8690 19;	19243	1.42	1.5E+00	6978492 NT	L	Rattus norvegicus 5 - Lipoxygenase (Alox5), mRNA
Ш				7661685 NT	7	Homo sapiens DKFZP586M0122 protein (DKFZP586M0122), mRNA
	10014 19809		1.4E+00	31685	Z	Homo sapiens DKFZP586M0122 protein (DKFZP586M0122), mRNA
2231 12	12118	0.85	1.4E+00	AF053357.1	ΡĀ	Helicobacter pylori glutamine synthetase (ginA) gene, complete cds
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Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acesslan No.	Top Hit Database Source	Top Hit Descriptor
2286			9:38	1.4E+00	U67922.1	NT	Ovis aries prion protein gene, complete cds
2834	12501	22394	1.63	1.4E+00	X74463.1	INI	Human papillomavirus type 7 genomic DNA
2746	12608	22500	3.29	1.4E+00	AF064584.2	TN	Fugu rubripes neurofibromatosis type 1 (NF1), A-kinase anchor protein (AKAP84), BAW protein (BAW), and WSB1 protein (WSB1) genes, complete cds
2746		22501	3.29	1.4E+00	AF064564.2	۲	Fugu rubripes neurofibromatosis type 1 (NF1), A-kinase anchor protein (AKAP84), BAW protein (BAW), and WSB1 protein (WSB1) genes, complete cds
3034		22765	1.01	1.4E+00	AE002324.2	NT	Chlamydia muridarum, section 55 of 85 of the complete genome
3034	12962	22756	1.01	1.4E+00	AE002324.2	NT	Chlamydia muridarum, section 55 of 85 of the complete genome
3291	13213		0.87	1.4E+00	5453733 NT	TN	Homo sapiens Mad4 homolog (MAD4) mRNA
4162	14062	23835	1.09	1.4E+00	AW800455.1	EST_HUMAN	CM0-NN1005-140300-288-h08 NN1008 Homo saplens cDNA
4162	14062	23836	1.09	1.4E+00	AW900455.1	EST_HUMAN	CM0-NN1005-140300-286-h06 NN1005 Homo sapiens cDNA
4482	14376		1.53	1.4E+00	BF681547.1	EST_HUMAN	602166687F1 NIH_MGC_83 Hamo saplens cDNA clone IMAGE:4297558 5'
5137	15004		8.0	1.4E+00	Q07869	SWISSPROT	PEROXISOME PROLIFERATOR ACTIVATED RECEPTOR ALPHA (PPAR-ALPHA)
5301	16222	26026	1.61	1.4E+00	AW054976.1	EST_HUMAN	wt45g07.x1 NCI_CGAP_Pan1 Homo sapiens cDNA clone IMAGE:2510460 3'
5394			5.17	1.4E+00	AB032983.1	ΝT	Homo sepiens mRNA for KIAA1157 protein, partial cds
5796	15702	25812	539	1.4E+00	013472	SWISSPROT	DNA TOPOISOMERASE III ALPHA
5802	19765		6'4	1.4E+00	AB020712.1	TN	Homo sapiens mRNA for KIAA0905 protein, complete cds
5860	15768	25884	2.71	1.4E+00	Q92777	SWISSPROT	SYNAPSIN II
5860	15766	25885	2.71	1.4E+00	Q92777	SWISSPROT	SYNAPSIN II
6318	16181	28341	1.86	1.4E+00	AJ133269.1	FX	Homo sapiens caveolin-1/-2 locus, Conüg1, D7Si322, genes CAV2 (exons 1, 2a, and 2b), CAV1 (exons 1 and 2)
7107			5.21	1.4E+00	AJ271735.1	LN	Homo sapiens Xq pseudoautosomal region; segment 1/2
7256	17133	27325	1.88	1.4E+00	R20459.1	EST_HUMAN	yg33f12.r1 Soares infant brain 1NIB Homo sapiens cDNA clone IMAGE:34345 5'
7303	17179	27381.	3.37	1.4E+00	BE084667.1	EST_HUMAN	RC1-BT0313-301299-012-105 BT0313 Homo saptens cDNA
8420	18294	28548	1.94	1.4E+00	AA195528.1	EST HUMAN	z/36e/9.r.1 Soares_NhHMPu_S1 Homo sapiens cDNA clone IMAGE:065512 5' similar to contains element MER22 repetitive element:
8556	Ι.	28695		1.4E+00	AB006682.1	N	Homo sapiens APECED mRNA for AIRE-1, complete cds
8709				1.4E+00	BE962107.2	EST HUMAN	601655184R1 NIH_MGC_65 Homo sapiens cDNA clone IMAGE:3846805 3'
8709				1.4E+00	BE962107.2	EST_HUMAN	601655184R1 NIH_MGC_65 Homo saplens cDNA clone IMAGE:3845805 3'
8727	18583	28867	2.68	1.4E+00	U30790.1	NT	Pneumocystis carinii f. sp. ratti guanine nucleotice binding protein alpha subunit (pcg1) gene, complete cds
8727	18583	28868	2.68	1.4E+00	U30790.1	LN	Pneumocystis carlnii f. sp. ratti guanine nucleotide binding protein alpha subunit (pcg1) gene, complete cds
9221	19604		1.34	1.4E+00	AL161500.2	LN	Arabidopsis thallana DNA chromosome 4, contig fragment No. 12

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פוופים דילון בספס ווין נפשו	Top Hit Descriptor	M.mucedo gene encoding 4-Dihydromethyl-trisporate dehydrogenase	Centharellus sp. partial 25S rRNA gene, isolate Tibet	Homo saplens putative psihHbA pseudogene for hair keratin, exons 2 to 7	Homo sapiens zinc finger protein 157 (HZF22) (ZNF157) mRNA	Homo sapiens zinc finger protein 157 (HZF22) (ZNF157) mRNA	Cotx lacryma-jobl dihydrodipicolinate synthase (dcpA) gene, complete cds	Chlamydia muridarum, section 66 of 85 of the corriplete genome	Opprinus carpio MRPb and MASPb genes for mannose-binding lectin-associated serine protease (MASP)	G01661233R1 NIH MGC 72 Homo saniens cDNA clone IMAGE:3915045 3'	Mus muscalus alpha-spectrin 1 arythroid (Spirat) mRNA	Control to be described to the control of the contr	rugu iudi pes ganinia-aniniooutynic ado receptor beta suounit gere, paruar cus, soku erynnocyte memorana protein (P55), synaptic vesicle-associated integral membrane protein (VAMP-1), procollagan C-proteinase	enhancer protein (PCOLCE) genes, complete c>	Mus musculus alpha-spectrin 1, erythroid (Spna1), mRNA	Piasmodium reichenowi partial 83/AMA-1 gene for apical membrane antigen 1	Piasmodium reichenowi partial 83/AMA-1 gene for apical membrane antigen 1	PMo-CT0289-291199-004-f08 CT0289 Homo sapiens cDNA	PM0-CT0289-291199-004-f08 CT0289 Homo saplens cDNA	D.melanogaster no-on-transient A gene product, complete ods	Sus scrafe plp gene	601657145R1 NIH_MGC_67 Homo saplens cDNA clone IMAGE:3866195 3'	Homo sepiens GL004 protein (GL004), mRNA	Homo sapiens heparan glucosaminyi N-deacetylase/N-sulfotransferase-2 gone, complete cds	Salba phr-1 mRNA for photolyase	S.alba phr-1 mRNA for photolyase	LYSOSOMAL ALPHA-MANNOSIDASE PRECURSOR (MANNOSIDASE, ALPHA B) (LYSOSOMAL ACID	ALPHA-MANNOSIDASE) (LAMAN)	wo85a07.x1 NCI_CGAP_Kid11 Homo saplens cDNA clone IMAGE:2462100 3'	601657145R1 NIH_MGC_67 Homo saplens cDNA clone IMAGE:3866195 3'	Vibrio cholerae chromosome II, section 49 of 93 of the complete chromosome	Campylobacter Jejuni kanamycin phosphotransferase (aphA-7) gene, complete cds	DIHYDROPYRIMIDINASE (DHPASE) (HYDANTOINASE) (DHP)	MRNA 3'-END PROCESSING PROTEIN RNA15
פון ווסעיין פוני	Top Hit Database Source	IN	IN	NT	TN	ᅜ	M	. LN	ţ	EST HIMAN	Ę		•	Ę	TN	IN	LN.	EST_HUMAN	EST_HUMAN	Ę	NT	EST_HUMAN	NT	TN	NT	IN		SWISSPROT		EST_HUMAN	ᅜ	NT	SWISSPROT	SWISSPROT
	Top Hit Acession No.	Z73840.1	2.1	Y19213.1	4507898 NT	4507898 NT	U61730.2	AE002338.2		AD030447.1				AF016494.1	6755621 NT	AJ252087.1	AJ252087.1	AW362834.1	AW362834.1	M33496.1	AJ009912.1	BE963379.2	10247	AF042084.1	X72019.1	X72019.1		000754		BE963379.2	AE004392.1	M28953.1	Q14117	P25289
		1.3E+00	1.3E+00	1.3E+00	1.3E+00	1.3E+00	1.3E+00		_	1.3E+00				1.3E+00	1.3E+00	1.3E+00	1.3E+00	1.3E+00	1.3E+00	1.3E+00		_	1.3E+00			1.3E+00			_		1.3E+00	1.3E+00	1.3E+00	1.3E+00
	Expression Signal	1.67	2.74	18.6	10.8	10.9	1.62	2.09	Č	2 10	1 43			1.03	0.82	86.0	96.0	7.70	7.78	1.33	1.25	3.06	1.55	5.44	2.31	2.31		1.45	1.27	4.9	1.6	1.39	4.3	2.34
	ORF SEQ ID NO:		20659			21034			-		22832	١		23258	22832		24608	25674		25889	26901	26988			27560	27561				1922	27961			28398
	- ₀	10498	10810	11028	1		11241	11497		12378	1			13463	12835	14839	14839	15576	15576	15770	16707	16795	16931	17350	17357	17357				17525	17718	17724		18157
		899	884	1113	1275	1275	1334	1593	27,00	2503	S S S			3547	4502	4964	4964	2999	5995	5864	6828	6917	7054	7480	7487	7487		7574	7620	7675	7866	7874	8078	8277

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Probe EX SEQ ID SEC NO: N	Exon ORF SEQ SEQ ID ID NO: NO:	EQ Expression O: Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
8299 18	18178 28	28424 2	2.23 1.3E+00	Z18892.2	IN	Mus musculus desmin gene
	18510	1.	1.81 1.3E+00	AW274791.1	EST_HUMAN	xp09e03.x1 NCI_CGAP_HN9 Homo sapiens cDNA clone IMAGE:2739868 3'
8826 18	18639 28	28923	2.82 1.3E+00	D42042.1	IN	Human mRNA for KIAA0085 gene, partial cds
8895 18	18705 28	28999 2.	2.71 1.3E+00	Z98682.1	N _T	Bacilius subtilis genomic DNA 23.9kB fragment
9362 18	19033	2	2.35 1.3E+00	AF187873.1	NT	Cavia porcellus inwardly-rectifying potassium channel Kir2.2 (KCNJ12) gene, complete cds
9533 16	19139 25	25263 3.	3.24 1.3E+00	BF348043.1	EST_HUMAN	602023185F1 NCI_CGAP_Bm67 Homo saplens cDNA clone IMAGE:4158452 5'
9544 16	19499	2.	2.68 1.3E+00	P33464	SWISSPROT	E1 GLYCOPROTEIN PRECURSOR (MATRIX GLYCOPROTEIN) (MEMBRANE GLYCOPROTEIN)
	10571 20:	20384 8.	8.14 1.2E+00	AA676246.1	EST_HUMAN	zi22d08.s1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:431535 3'
	10735 20	20578 1.	L	P05228	SWISSPROT	HISTIDINE-RICH PROTEIN PRECURSOR (CLONE PFHRP-III)
L	10735 20	20579 1.	1.33 1.2E+00	P05228	SWISSPROT	HISTIDINE-RICH PROTEIN PRECURSOR (CLONE PFHRP-III)
808 10	10735 20	20580 1.	1.33 1.2E+00	P05228	SWISSPROT	HISTIDINE-RICH PROTEIN PRECURSOR (CLONE PFHRP-III)
860 10	10786	2	2.19 1.2E+00		IN	Homo saplens hypothetical protein PRO3077 (PRO3077), mRNA
1145 11	11058 20		5.53 1.2E+00	AF080245.2	ĹΝ	Elasis oleifera sesquiterpene synthase mRNA, complete cds
1187 11	11097 20	20943 1.	1.77 1.2E+00	AJ252242.1	TN	pea seed-borne mosaic virus complete genome
1187 1	11097 20		1.77 1.2E+00	AJ252242.1	M	pea seed-borne mosaic virus complete genome
1963 1	11857 21	21746 1.	1.05 1.2E+00	AF140631.1	NT	Homo sapiens G-protein coupled receptor 14 (GPR14) gene, complete cds
	12999 22	22789 0.	0.95 1.2E+00	AB020881.1	L	Homo sapiens mRNA for KIAA0874 protein, partial cds
				AL161563.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 63
3128 13	13053 22	22852 5.	5.41 1.2E+00	AL161563.2	LN	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 63
3249 1	13172		2.9 1.2E+00	P54910	SWISSPROT	CONJUGAL TRANSFER PROTEIN TRBE PRECURSOR
3311 13	13232 23	23038 0.	0.78 1.2E+00	AF188740.1	NT	Homo sapiens LHX3 gene, intron 2
	13233	0	0.89 1.2E+00	M81779.1	Ę	G.gallus T-cadherin mRNA, complete cds
	13564 23		7.28 1.2E+00	U75902.1	NT	Mus musculus subtilisin-like serine protease LPC (PC7) gene, exons 1 to 9, partial cds
			1.49 1.2E+00	BF373570.1	EST_HUMAN	MR0-FT0175-050900-203-g06_1 FT0175 Homo sapiens cDNA
4201 1	13232 23	23038 1.	1.09 1.2E+00	AF188740.1	Ŋ	Homo sapiens LHX3 gene, intron 2
	14196 23	23981 1.	1.39 1.2E+00	0 6980951 NT	IN	Rattus norvegicus Glycine receptor alpha 2 subunit (glycine receptor, neonatal) (Glra2), mRNA
4371 1	14267	1	1.64 1.2E+00	M87060.1	IN	Rattus rattus cardiac AE3 gene, exons 1-23
4422 1	14316 24	24102	1.28 1.2E+00	AL161509.2	IN	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 21
4460 1	14354 24	24145 1.	1.82 1.2E+00	AF158495.1	TN	Homo sapiens post-synaptic density 95 (DLG4) gene, complete cds
4485 1	14379	5	5.44 1.2E+00	Y09200.1	INT	T.pinnatum chloroplast rbcL gene, partial
4584 1	13233	0	0.94 1.2E+00	M81779.1	IN	G.gallus T-cadherin mRNA, complete cds.
			1.96 1.2E+00	AW813276.1	EST_HUMAN	MR3-ST0191-140200-013-c05 ST0191 Homo sapiens cDNA
				X74885.1		D.hydel ayf repeat cluster DNA, fragment D
5769 1	15676 25	25783 3.	3.28 1.2E+00	D BE003113.1	EST_HUMAN	QV4-BN0090-270400-190-e03 BN0090 Hamo sapiens cDNA

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	riptor			le 1322374 3'	NA					NA	ructokinase beta subunit	d clone hm01a01		NA	NA	complete cds	mplete cds				NA	s cDNA clone IMAGE:3074834 3'			mRNA	mRNA	A clone IMAGE:2359461 3' similar to	Jr. 1,	9	e		mRNA		nRNA, complete cds	
oo Expressed in France	Top Hit Descriptor	C.glutamicum pta gene and ackA gene	C.glutamicum pta gene and ackA gene	ah84g12.s1 Soares_testis_NHT Homo sapiens cl)NA clone 1322374 3'	MR3-ST0191-140200-013-c05 ST0191 Homo saplens cDNA	Mus musculus DSPP gene	AV734585 cdA Homo sapiens cDNA clone cdAAFH03 5	Lilactis pyrD and pyrF genes	Homo sepiens mRNA for KIAA1204 protein, partial cds	MR2-CT0222-201099-001-607 CT0222 Homo sapiens cDNA	R.communis gene for pyrophosphate-dependent phosphofructokinase beta subunit	HUMHM01A01 Liver HepG2 cell line. Homo sapiens cDNA clone hm01a0	H.saplens ENO3 gene for muscle specific enolase	PM0-ST0264-161199-001-d01 ST0264 Homo sapiens cDNA	PM1-HT0422-160200-007-910 HT0422 Homo saplens cDNA	Rattus norvegicus synapse-associated protein 10,2 mRNA, complete cds	Mus musculus 60 kDa ribonucleoprotein SSA/Ro gene, complete cds	Homo saplens chromosome 21 segment HS21C003	Bacillus halodurans genomic DNA, section 9/14	Human mRNA for KIAA0227 gene, partial cds	QV0-BN0042-170300-163-912 BN0042 Homo seplens cDNA	UI-HF-BR0p-ak-f-02-0-UI.s1 NIH_MGC_52 Homo saplens cDNA clone IMAGE:3074834 3'	Homo sapiens chromosome 21 segment HS21C013	Homo sapiens chromosome 21 segment HS21C013	Homo saplens hypothetical protein FLJ10749 (FLJ10749), mRNA	Homo sapiens hypothetical protein FLJ11280 (FLJ11280), mRNA	wf54h11x1 Soares_NFL_T_GBC_S1 Home sapiens cDNA clone IMAGE:2359461 3' similar to	SWITSSI HUMAN GIZSSO POS-BINDING PROJECT SSEPTI	Ayella fastidiosa, section 32 of 229 of the complete genome	Xylella fastidiosa, section 32 of 229 of the complete genome	Homo sepiens calpain 9 (nCL-4) (CAPN9) mRNA.	Homo sapiens hypothetical protein FLJ10749 (FLJ10749), mRNA	R.unicornis complete mitochondrial genome	Drosophila melanogaster cytoplasmic dynein heavy chain mRNA, complete cds	Mrican swine fever virus complete denome
301 1 100 P	Top Hit Database Source	NT	M	EST_HUMAN	EST_HUMAN	Ŋ	EST_HUMAN	NT	NT	EST_HUMAN		EST_HUMAN	INT	EST_HUMAN	EST_HUMAN	IN	NT	MT	NT	IN	EST_HUMAN	EST_HUMAN	NT	NT	F	ŊŢ		ESI TOMAN	IN	TN	뒫	INT	NT	Ę	5
	Top Hit Acessian No.	X89084.1	X89084.1	AA759254.1	4W813276.1	AJ002141.1	AV734585.1	X74207.1		AW377210.1	232850.1	D11745.1	X56832.1	AW817817.1	BE160761.1	J50147.1	AF065398.1	AL163203.2	AP001515.1	D86980.1	AW895393.1	AW575889.1	AL163213.2	AL163213.2	8922641 NT	8922973 NT	7 0000001	AIBUBSBO.1	AE003886.1	AE003886.1	5729767 NT	8922841 NT	5835331 NT	L23195.1	110/00 4
	Most Similar (Top) Hit BLAST E Value	1.2E+00	1.2E+00 >	1.2E+00/	1.2E+00	-	1.2E+00/			1.2E+00	•	_	1.2E+00 >	1		-		1.2E+00	1.2E+00	1.1E+00	1.1E+00/	1.1E+00 /	_		1.1E+00	1.1E+00				_			1.1E+00	1.1E+00	4 4E+00
	Expression Signal	1.87	1.87	32.83	1.82	2.55	د. و:	2.4	3.45	1.74	3.08	1.8	3.28	. 2.03	23.47	4.87	2.5	15.4	1.49	1.13	1.39	0.92	6.61	9.61	0.83	1.26		0.99	1.18	1.16	17.81	0.99	5.89	1.18	202
	ORF SEQ ID NO:	25815	25816	25841	. 25949	26121	26392	26544	27055		27480	27601	27786	28826		28145	28787	25087		20217	21486	21628	23009	23010	23165		,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	23244	23357	23358	23378	23565			5257C
	Exon SEQ ID NO:	15704	15704	15729	15823	15986	19463	16367	16861	17095	17274	17391	17561	18542	18576	17901	18877	19572	18025	10400	11627	11753		13209	13358	13426	-11,00	1				13773	14022		14800
	Probe SEQ ID NO:	5798	5798	5823	5918	6139	6369	8208	6984	7218	7407	7540	7711	8653	8689	8752	9114	9330	9349	458	1726	1857	3288	3288	3441	3510	7 0	3531	3657	3657	3678	3862	4122	4860	4024

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Г						_	_								_							_	7	7		-		7			\neg
	Top Hit Descriptor	E.faecalis pbp5 gene	Rattus norvegicus Aquaporin 4 (Aqp4), mRNA	601652776R1 NIH_MGC_58 Hamo sapiens cDNA clone IMAGE:3826835 3'	qd85c03.x1 Soares_testis_NHT Hamo sapiens clJNA clone IMAGE:1736260 3'	Herpes simplex virus type 1 (strain KOS) UL41 gene	Herpes simplex virus type 1 (strain KOS) UL41 gene	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 84	602082582F1 NIH_MGC_81 Homo sapiens cDNA clone IMAGE: 4246628 5'	Homo saplens mRNA for KIAA0934 protein, partial cds	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 27	Mus musculus guanine nucleotide binding protein (G protein), gamma 3 subunit (Gng3), mRNA	Homo saplens KIAA0626 gene product (KIAA0626), mRNA	Klebsormidium fluitans cytochrome c oxidase subunit 2 (cox2) gene, mitochondrial gene encoding mitochondrial matein partial cyte.	Homo sapiens hypothetical protein FLJ11280 (FLJ11280), mRNA	Petroselinum crispum cytosolic glucose-6-phosphate dehydrogenase 1 (cG6PDH1) mRNA, complete ods	Petroselinum crispum cytosolic glucose-6-phosphate dehydrogenase 1 (cG6PDH1) mRNA, complete cds	wf76e11.x1 Scares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2361548 3'	LOW TEMPERATURE ESSENTIAL PROTEIN	Taenia solium immunogenic protein Ts76 mRNA, partial cds	Dictycstellum discoldeum isopentenyl pyrophosphate isomerase (Dipl) mRNA, complete cds	Xenopus laevis rhodopsin gene, complete cds	Cavia cobaya mRNA for serine/threoine kinase, complete cds	Marchantia polymorpha genes for 26S rRNA, 5S rRNA, 18S rRNA, 5.8S rRNA and 26S rRNA	Girardia tigrina mRNA for homeodomain transcription factor (so gene)	Homo saplens chromosome 21 segment HS21C018	Aedes aegypti mucin-like protein MUC1 mRNA, complete cds	V.carteri Algal-CAM mRNA	Plautia stall intestine virus RNA for nonstructural polyprotein, capsid protein precursor, complete cds	DNA GYRASE SUBUNIT B	DNA GYRASE SUBUNIT B
	Top Hit Database Source	N	N	EST_HUMAN	EST_HUMAN	Z	Ν	F	EST_HUMAN	Ę	뒫	NT	ΝŢ	Ę	5 5	Ę	Ę	EST_HUMAN	SWISSPROT	NT	NT	NT	M	TN	IN	NT	NT	Ν	Ā	SWISSPROT	SWISSPROT
	Top Hit Acession No.	X78425.1	FN 0538580	BE960184.1	Al138582.1	Z7238.1	Z72338.1	AL161588.2	BF693996.1	AB023151.1	AL161515.2	6754021	11067364 NT	A Enesono 4	8922973 NT	AF012862.1	AF012862.1	AI809699.1	P07868	AF216696.1	AF234169.1	U23808.1	D88425.1	AB021684.1	AJ251660.1	AL163218.2	AF125984.1	X80416.1	AB006531.1	P48355	P48355
	Most Similar (Top) Hit BLAST E Value	1.1E+00	1.1E+00	1.1E+00	1.1E+00		-	1.1E+00	1.1E+00	1.1E+00	1.1E+00	1.1E+00	1.1E+00	4 45,00	_		-	1.1E+00	1.1E+00	1.1E+00	1.1E+00	1.0E+00	1.0E+00			1.0E+00	1.0E+00	1.0E+00	1.0E+00		1.0E+00
	Expression Signal	1.18	1.57	12.04	1.29	1.93	1.93	7.53	3.22	1.65	4.39	20.08	265		5.73	4.41	4.41	4.73	4	2.13	1.54	3.49	1.14	2.16	1.14	4.51	0.95	1.04	1.32	1.4	1.4
	ORF SEQ ID NO:	24638	24945	25426	25440		L	26468	26789	27733	27791	27815	28182			28581	28582			25310			19904			20413			21492		
	Exon SEQ ID NO:	14874	15172	15370	15380		L	16296	16608	17508	17565	17594	17934	4 7004	1	<u></u>	<u> </u>	18551	18999	19056	19570	1001	10089	10358	10502	10595	10597	12692	11623	1	1 1
	Probe SEQ ID NO:	4999	5249	5449	6460	6422	6422	6435	6728	7658	7716	7744	8043	8	8447	8450	8420	8662	8300	9402	9523	92	108	412	562	661	ශි	1364	1722	2436	2435

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Table 4
Single Exon Probes Expressed in Heart

					"		
Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	. Top Hit Descriptor
2651	12518	22408	1.09	1.05+00	AF131205.1	Z	Mus musculus Serf1 protein (Serf1), survival of motor neuron protein (Smn), neuronal apoptosis inhibitory protein-rs6 (Naip-าร6), and neuronal apoptosis inhibitory protein-rs3 (Naip-rs3) genes, complete cds
2846	12774	22661	3.48	1.0E+00	P24008	SWISSPROT	3-OXO-5-ALPHA-STEROID 4-DEHYDROGENASE 1 (STEROID 5-ALPHA-REDUCTASE 1) (SR TYPE 1)
2848	12774	22562	3.48	1.0E+00	P24008	SWISSPROT	3-OXO-5-ALPHA-STEROID 4-DEHYDROGENASE 1 (STEROID 5-ALPHA-REDUCTASE 1) (SR TYPE 1)
2938	12865		0.89	1.0E+00	014226	SWISSPROT	HYPOTHETICAL 67.9 KD PROTEIN C6F12.08C IN CHROMOSOME I
3162	13087	22891	1.17	1.0E+00		EST_HUMAN	ar26g08.s1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA clone IMAGE:1032830 3' similar to WP:C42D8.3 CE04204 :contains element MER22 MER22 repetitive element ;
3327	13247		0.81	1.0E+00	AF222761.1	NT	Rattus norvegicus neuromedin U precursor (NmU) gene, exons 5 and 6
3548	10077		1.15		U23808.1	TN	Xenopus laevis rhodopsin gene, complete cds
3627	13541	23328	1.44	1.0E+00	.1	TN	Agaricus bisporus mRNA for tyrosinase
3978	13885	23660	0.86	1.0E+00	AF2Z3391.1	NT	Homo sapiens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively spliced
4178	14078		0.79	1.0E+00	8922245 NT	LN LN	Homo sapiens hypothetical protein FLJ10139 (FLJ10139), mRNA
4815	14698		0.88	1.0E+00	U75741.1	NT	Taenia ovis 45W antigen (ToW4) gene, complete cds
4951	14828		0.88	1.0E+00	D10852.1	TN	Rattus norvegicus mRNA for N-acetylglucosaminyltransferase III, complete cds
5139	15008	24777	0.87	1.0E+00	AJ223978.1	N	Bacillus subtilis 42.7kB DNA fragment from yvsA to yvqA
5228	15152	24919		1.0E+00		NT	Hordeum vulgare gene encoding cystalne proteinase
5568	15484	25557	4.31	1.0E+00	AF248054.1	TN	Bos teurus micromolar calcium activated neutral protease 1 (CAPN1) gene, exons 11-20, and partial cds
5568	15484	25558	4.31	1.0E+00	AF248054.1	LΝ	Bos taurus micromolar calcium activated neutral protease 1 (CAPN1) gene, exons 11-20, and partial cds
5719	15626	25729		1.0E+00	P04501	SWISSPROT	FIBER PROTEIN
5720	15627	25730	1.39	1.0E+00	82.1	EST_HUMAN	ULH-BIS-alx-d-09-0-UI.s1 NCI_CGAP_Sub5 Homo sapiens cDNA clone IMAGE:3068969 3'
5908	15814	25939	2.21	1.0E+00	U75902.1	NT.	Mus musculus subtilisin-like serine protease LPC (PC7) gene, exons 1 to 9, partial cds
6417	16279		8.43	1.0E+00	P20273	SWISSPROT	B-CELL RECEPTOR CD22 PRECURSOR (LEU-14) (B-LYMPHOCYTE CELL ADHESION MOLECULE) (BL-CAM)
6537	16395	26574	1.34	1.0E+00	AF192531.1	Į.	Homo sapiens endothelin-converting enzyme 2 (ECE2) mRNA, complete cds
8548	16404	28583	6:88	1.0E+00	AA776191.1	EST_HUMAN	ac79b08.s1 Stratagene lung (#537210) Homo sapiens cDNA clone IMAGE:868791 3'
6841		26713			BE868267.1	EST_HUMAN	601443950F1 NIH_MGC_65 Homo sapiens cDNA clone IMAGE:3848005 5'
6841		26714		1.0E+00	BE868267.1	EST_HUMAN	601443950F1 NIH_MGC_65 Homo seplens cDNA clone IMAGE:3848005 5'
6734	14828		1.38	1.0E+00	D10852.1	Ę	Rattus norvegicus mRNA for N-acetylglucoseminyltransferase III, complete cds

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Exon ORF SEQ Expression (Top) Hit T Signal BLASTE Value	Most Similar Expression (Top) Hit Signal BLAST E Value	Most Similar (Top) Hit BLAST E Value		<u> </u>	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
16740 26932 2.7 1.0E+00 002	26932 2.7 1.0E+00	2.7 1.0E+00	1.0E+00		002207	SWISSPROT	PEROXISOMAL HYDRATASE-DEHYDROGENASE-EPIMERASE (HDE) (MULTIFUNCTIONAL BETA- OXIDATION PROTEIN) (MFP) [INCLUDES: 2-ËNOYL-COA HYDRATASE ; D-3-HYDROXYACYL COA DEHYDROGENASE]
16740 28933 2.7 1.0E+00 Q02207	26933 2.7 1.0E+00(2.7 1.0E+00(1.0E+00		20	SWISSPROT	PEROXISOMAL HYDRATASE-DEHYDROGENASE-EPIMERASE (HDE) (MULTIFUNCTIONAL BETA- OXIDATION PROTEIN) (MFP) [INCLUDES: 2-ENOYL-COA HYDRATASE ; D-3-HYDROXYACYL COA DEHYDROGENASE]
19466 2.01 1.0E+00 BE14	2.01 1.0E+00	1.0E+00	3	3	3E147331.1	EST_HUMAN	RC1-HT0229-181099-011-e06 HT0229 Homo sapiens cDNA
27133 1.9 1.0E+00	27133 1.9 1.0E+00	1.9 1.0E+00	1.0E+00	ł	M38427.1	NT	Human immunodeficiency virus type 1 (HIV-1), isolate SF33,
1.76 1.0E+00 B	27405 1.76 1.0E+00 B	1.76 1.0E+00 B	1.0E+00 B	8	E907592.1	EST_HUMAN	601497581F1 NIH_MGC_70 Homo saplens cDNA clone IMAGE:3899421 5'
27526	27526 1.28	1.28			6753429 NT	NT	Mus musculus chloride channel calcium activated 1 (Cica1), mRNA
27527 1.28 1.0E+00	27527 1.28 1.0E+00	1.28 1.0E+00	1.0E+00		6753429 NT	NT	Mus musculus chloride channel calcium activated 1 (Cloa1), mRNA
27607 2.03 1.0E+00/	27607 2.03 1.0E+00	2.03 1.0E+00	1.0E+00/		AV689554.1	EST_HUMAN	AV689554 GKC Homo seplens cDNA clone GKCCYA11 5'
17397 27609 1.2 1.0E+00 U44952.1	1.2 1.0E+00 L	1.2 1.0E+00 L	1.0E+00	_	12.1	NT	Xenopus laevis zona pelludda C glycoprotein precursor (xIZPC) mRNA, complete cds
17397 27610 1.2 1.0E+00 U44952.1	27610 1.2 1.0E+00 L	1.2 1.0E+00	1.0E+00	1	21	NT	Xenopus laevis zona pellucida C glycoprotein precursor (xZPC) mRNA, complete cds
2.83	28005 2.83 1.0E+00	2.83 1.0E+00	1.0E+00 /	1	825.1	EST_HUMAN	AV758825 BM Homo sapiens oDNA clone BMFAW C04 5
17.08 1.0E+00[/	100+30.1 80.71 1.0E+00	17.08 1.0E+00	1.0E+00	1	982.1	EST_HUMAN	zh94a02.r1 Soares_fetal_liver_spleen_1NFLS_S1 Homo saplens cDNA clone IMAGE:428906 5'
28076 17.08 1.0E+00/	28076 17.08 1.0E+00/	17.08 1.0E+00/	1.0E+00/	•	382.1	EST_HUMAN	zh94a02.r1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:428906 5'
18928 2.05 1.0E+00 P15306	2.05 1.0E+00 F	1.0E+00	1.0E+00	_	3	SWISSPROT	THROMBOMODULIN PRECURSOR (FETOMODULIN) (TM)
1.0E+00	1.51 1.0E+00	1.0E+00	1.0E+00	1	5184.1	EST_HUMAN	EST388293 MAGE resequences, MAGN Homo saplens cDNA
1.67 1.0E+00	1.67 1.0E+00	1.0E+00 /	1.0E+00 /	•	50.1	ΝΤ	Homo sapiens mRNA for KIAA1517 protein, partial cds
0.84 9.9E-01	21315 0.84 9.9E-01	0.84 9.9E-01	9.9E-01	_	455.1	NT	Drosophila melanogaster regulator of G-protein signalling LOCO III mRNA, complete cds
21316 0.84 9.9E-01 /	21316 0.84 9.9E-01	0.84 9.9E-01	9.9E-01	•	455.1	NT	Drosophila melanogaster regulator of G-protein signalling LOCO III mRNA, complete cds
22360 0.9 9.9E-01 /	22360 0.9 9.9E-01 /	0.9 9.9E-01	9.9E-01	_	4L163302.2	NT	Homo sapiens chromosome 21 segment HS21C/102
1.18 9.9E-01/	1.18 9.9E-01	1.18 9.9E-01/	9.9E-01 /	•	4F174585.1	NT	Apple mosaic virus RNA 2 putative polymerase gene, complete cds
	25441 8.22 8.9E-01	8.22 8.9E-01	9.9E-01	-	57	SWISSPROT	SERINE/THREONINE PROTEIN KINASE MINIBRAIN
	1.38 9.9E-01	10-36.6	10-36.6		J65667.1	TN	Lycopersicon esculentum putative Mi1 copy 1 nematode-resistance gene
	2.43 9.9E-01	9.9E-01	9.9E-01		42	SWISSPROT	B2 BRADYKININ RECEPTOR (BK-2 RECEPTOR)
-	28237 2.92 9.9E-01	2.92 9.9E-01	9.9E-01	-	1,005029.1	TN	Danto reno mRNA for Eph-like receptor tyrosine kinase rtk8
10454 20265 2.32 9.8E-01 P22567	20265 2.32 9.8E-01	2.32 9.8E-01	9.8E-01	_	22	SWISSPROT	AMINO-ACID ACETYLTRANSFERASE (N. ACETYLGLUTAMATE SYNTHASE) (AGS) (NAGS)
9.8E-01	23429 0.82 9.8E-01	0.82 9.8E-01	9.8E-01		3E957439.2	EST_HUMAN	601653583R2 NIH_MGC_55 Homo saplens cDNA clone IMAGE:3838461 3'
9.8E-01	23430 0.82 9.8E-01	0.82 9.8E-01	9.8E-01		3E957439.2	EST_HUMAN	601653583R2 NIH_MGC_55 Homo sapiens cDNA clone IMAGE:3838461 3'
16137 26292 4.61 9.8E-01 AJ30	26292 4.61 9.8E-01	4.61 9.8E-01	9.8E-01		1302158.1	TN	Enterobacteriaceae sp. JM983 partial groES gens for GroES-like protein and partial groEL gene for GroEL- like protein, isolate JM983

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Table 4
Single Exon Probes Expressed in Heart

Top Hit Descriptor	Enterobacteriaceae sp. JM983 partial groES gene for GroES-like protein and partial groEL gene for GroEL- like protein, isolate JM983	601110258F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3350750 5'	601110258F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3350750 5	Homo sapiens X28 region near ALD locus containing dual specificity phosphatase 9 (DUSP9), ribosomal protein L18a (RPL18a), Ca2+/Calmodulin-dependent protein kinase I (CAMKI), creatine transporter (CRTR), CDM protein (CDM) adrenoleukodystrophy protein >	we62e04.x1 Soares_thymus_NHFTh Homo sapiens cDNA clone IMAGE:2345694 3'	Drosophila melanogaster sodium channel protein (para) gene, exons 9,10,11,12 and optional segments b, c, d	and of parts cus. Triticum aestivum stribe rust resistance protein Yi10 (Yr10) gene, complete cds	Salmonella typhimurium adenine-methyltransferase (mod) and restriction endonuclease (res)	UI-H-BI4-aci-e-07-0-UI.s1 NCI_CGAP_Sub8 Homo sapiens cDNA clone IMAGE:3086140 3'	Botrytis cinerea strain T4 cDNA library under conditions of nitrogen deprivation	Bramus inermis putative cylosolic phosphoglucoriutase (pgm1) mRNA, complete cds	Bromus inermis putative cytosolic phosphoglucomutase (pgm1) mRNA, complete cds	PM2-UM0053-240300-005-f12 UM0053 Homo sapiens cDNA	Homo sapiens KIAA0914 gene product (KIAA0914), mRNA	Parvovirus B19 DNA, patient C, genome position 2448-2994	Parvovirus B19 DNA, patient C, genome position 2448-2994	P.falciparum complete gene map of plastid-like DINA (IR-A)	AV752605 NPD Homo sapiens cDNA clone NPDBAG06 5'	AV752605 NPD Homo septens cDNA clone NPDBAG08 6'	Sphyrna tiburo NADH dehydrogenase subunit 2 (NADH2) gene, mitochondrial gene encoding mitochondrial	Protest, per usi ous 60487583061 NIH MCC 24 Homo emises -DNA Acres IMACE 3058473 F	801875830E1 NIH MGC 21 Homo samions CDNA Acros NAACE 3058479 5	RC1-CT0295-241199-011-b02 CT0295 Home sablens cDNA	Bartonella clarridgelae RNA polymerase beta subunit (rpoB) gene, partial cds	Pimpinella brachycarpa zinc finger protein (ZFP1) mRNA, complete cds	601466703F1 NIH_MGC_67 Homb sapiens cDNA clone IMAGE:3869929 5'	Homo sapiens epidermal growth factor receptor (avian erythroblastic leukemia viral (v-erb-b) oncogene homolog) (EGFR), mRNA	Homo saplens phytanoyl-CoA hydroxylase (PHYH) gene, expn 5
Top Hit Database Source	<u> </u>	EST_HUMAN 6	EST_HUMAN 6	I d C	T HUMAN				T	F E			EST HUMAN F		H IN				EST_HUMAN A		I I MAN	Т	Т		P.	EST_HUMAN 6		
Top Hit Acession No.	AJ302158.1	BE258705.1	BE258705.1	U52111.2	AI660384.1	100746 4	AF149112.1	M90544.1	-	AL114281.1	AF197925.1	AF197925.1	AW799674.1	7662375 NT	Z70556.1	Z70556.1	X95275:1	AV752605.1	AV752605.1	100,400	DE002340 4	REGOZZAO 4	AW861102.1	AF165990.1	AF080595.1	BE781251.1	11419857 NT	AF242382.1
Most Similar (Top) Hit BLAST E Value	9.8E-01	9.8E-01	9.8E-01	9.8E-01	9.7E-01	11				9.7E-01	9.6E-01	9.6E-01	9.6E-01	9.6E-01	9.6E-01	9.6E-01			9.6E-01	10.0				9.4E-01	9.4E-01	9.4E-01	9.4E-01	
Expression Signal	4.61	4.43	4.43	1.62	1.01	202	1.88	1.74	4.62	1.39	6.8	6.8	1.3	0.92	3.46	3.46	1.37	4.15	4.15	62.4	36.1	7	1.17	3.37	1.67	1.64	1.43	1.11
ORF SEQ ID NO:			28488			aacac	27023	27025			24016	. 24017	24036	24729		25499			28913	000		23411	27336					
Exen SEQ ID NO:	16137	18239	18239	19061	1	18114		1	18382	19419			1	14953					18622	97907		ł	1	L	13106	19029	19564	11601
Probe SEQ ID NO:	6272	8362	8362	9408	5108	8778	6952	6954	8510	9366	4337	4337	4355	5083	5517	5517	6890	8808	8808	0020	3715	3718	7266	3165	3181	9354	8026	1700

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Тop Hit Descriptor	RC5-BT0503-271199-011-B01 BT0503 Homo sapiens cDNA	Bovine papiliomavirus type 2, complete genome	Bovine papillomavirus type 2, complete genome	Equus caballus microsatellite LEX013	Homo sapiens nuclear factor of kappa light polypeptide gene enhancer in B-cells 1 (NFKB1) gene, complete	SOO.	Spodoptera frugiperda methyteneterranydrotolate dehydrogenase mRNA, complete cds	oe09b03.s1 NCLCGAP_Ov2 Home sapiens cDNA clone IMAGE:1385357	Aedes triseriatus putative large subunit ribosomal protein rpL34 mRNA, complete cds	Homo sapiens chromosome Xq28 melanoma antigen family A2a (MAGEA2A), melanoma antigen family A12 (MAGEA12), melanoma antigen family A2b (MAGEA2B), melanoma antigen family A3 (MAGEA3), caltractin	(CALT), NAD(P)H dehydrogenase-like protein (NSDHL), and LI>	601441338T1 NIH_MGC_72 Homo sapiens cDNA clone IMAGE:3916184 3'	601461153F1 NIH_MGC_66 Homo sapiens cDNA clone IMAGE:3864681 5'	Mus musculus carbonic anhydrase 4 (Car4), mRŅA	Homo sapiens lysosomal apyrase-like protein 1 (LALP1), mRNA	7o58e06.x1 NCI_CGAP_Kid11 Homo sepiens cDNA clone IMAGE:3578219 3' eimiler to SW:NU5M_TRYBB Po4540 NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 5;	601820312F1 NIH_MGC_58 Homo sapiens cDNA clone IMAGE:4052018 5'	602154769F1 NIH_MGC_83 Homo sapiens cDNA clone IMAGE:4295896 5	Homo sapiens hypothetical protein FLJ20048 (FLJ20048), mRNA	AB200G8R Infant brain, LLNL array of Dr. M. Socires 1NIB Homo septens cDNA clone LLAB200G8 5"	AB200G8R Infant brain, LLNL array of Dr. M. Soeires 1NIB Homo sapiens cDNA clone LLAB200G8 6'	Rattus norvegicus mucin (MUC2) gene, partial cds	INTER-ALPHA-TRYPSIN INHIBITOR HEAVY CHAIN H3 PRECURSOR (ITI HEAVY CHAIN H3)	ob71g08.s1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:1336862 3'	Rattus norvegicus Rab3 GDP/GTP exchange protein mRNA, complete cds	Homo sapiens uncoupling protein-3 (UCP3) gene, complete cds	Homo sapiens DKFZP564M2423 protein (DKFZP564M2423), mRNA	Homo sapiens neurexin III-alpha gene, partial cds	Oryctolagus cuniculus Rad51 (RAD51) mRNA, complete cds	Xenopus laevis gene for aldolase, complete cds
Top Hit Database Source	EST_HUMAN	IN	NT	NT	Ŀ	Į.	Ę	EST_HUMAN	L		뉟	EST_HUMAN	EST_HUMAN	NT	ΙN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	EST_HUMAN	EST_HUMAN	۲	SWISSPROT	EST_HUMAN	TN	IN	NT	NT	NT	NT
Top Hit Acession No.	BE071172.1	9.3E-01 M20219.1	9.3E-01 M20219.1	9.3E-01 AF075615.1		9.3E-01 AFZ13884.1	ı	-	9.3E-01 AF271207.1		9.3E-01 U82671.2	9.2E-01 BE622702.1	BF037586.1	6671677 NT	11430963 NT	9.2E-01 BF593251.1	BF132402.1	BF680047.1	8923056 NT	9.1E-01 T26418.1	T26418.1	U68172.1	Q61704		U72995.1	AF060113.1	7661625 NT	AF099810.1	9.0E-01 AF017729.1	D38621.1
Most Similar (Top) Hit BLAST E Value	9.3E-01 B	9.3E-01	9.3E-01	9.3E-01	70.00	1	ł	ı			9.3E-01	9.2E-01	9.2E-01	9.2E-01	9.2E-01	9.2E-01	9.2E-01	9.2E-01 BF	9.1E-01	9.1E-01	9.1E-01	9.1E-01	9.1E-01 Q61704	9.1E-01	9.1E-01	9.1E-01	9.0E-01	9.0E-01 A	9.0E-01	9.0E-01
Expression Signal	1.13	0.82	0.82	1.54		1.62	3.84	1.88	1.56	i	1.38	3.04	4.23	1.38	3.63	1.73	1.85	2.01	1.43	78.0	0.97	0.86	2.92	15.99	2.2	30.05	0.89	2.37	0.82	1.45
ORF SEQ ID NO:		23630	23631				25465					22930	25649				29059	29101		22898			25950	26498				23959		
Exon SEQ ID NO:	12464	13857	13857	15022		L	_ 1	- 1	19325		19418	13125	15558	1	17740	17802	18767	1	11967	13094	13094		15824	16331	16409	19641	13095	14181		16250
Probe SEQ ID NO:	2594	3949	3949	5155		454	5482	6701	9822		9954	3201	5643	7618	7890	7952	0968	9006	2077	3169	3169	4310	5919	6472	6551	9446	3170	4282	4946	6388

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Single Exon Probes Expressed in Heart

		, -		_		-	_		_	_			_	_		_	_	_	_				_	_		_		_		
Top Hit Descriptor	Fugu rubripes neural cell adhesion molecule L1 homolog (L1-CAM) gene, complete cds; putative protein 1 (PUT1) gene, partial cds; mitosis-specific chromosome segregation protein SMC1 homolog (SMC1) gene, complete cds; and calcium channel alpha-1 subunits	Rabbit MHC fragment RLA-DF DNA	Xylella fastidiosa, section 90 of 229 of the complete genome	Chlamydophila pneumoniae AR39, section 21 of 94 of the complete genome	PUTATIVE F420-DEPENDENT NADP REDUCTASE	M.aeruginosa (HUB 5-2-4) DNA from plasmid PN/A1	Synechocystis sp. PCC6803 complete genome, 13/27, 1576593-1719643	Homo sepiens SOS1 (SOS1) gene, partial cds	Homo sapiens AT-binding transcription factor 1 (ATBF1), mRNA	Inn05f11.s1 NCI_CGAP_Pr4.1 Homo sapiens cDNA clone IMAGE:1076877	Pseudomonas aeruginosa topolsomerase (top), putative transcriptional regulatory protein OhbR (ohbR), ortho- haidbenzosta 1 2-diroxdenasa bata-ISP protein OhbA (ohbA), OhbC (ohbC), ortho-haidbenzosta 1 2-	dioxygenase alpha-ISP protein OhbB (ohbB), and put>	Pseudomonas aeruginosa PA01, section 524 of 529 of the complete genome	QV0-NN1021-100800-337-c03 NN1021 Hamo sapiens cDNA	601823684R1 NIH MGC_79 Homo sapiens cDNA clone IMAGE:4043564 3'	601823684R1 NIH_MGC_79 Homo saplens cDNA clone IMAGE:4043564 3'	AV661898 GLC Homo sapiens cDNA clone GLCGYG07 3'	Rat IGFII gene for insulin-like growth factor II	zd44e03.r1 Soares_fetal_heart_NbHH19W Homo sapiens cDNA clone IMAGE:343516 5	Homo sepiens cytochrome P450, subfamily XXVIIIA (steroid 27-hydroxylase, cerebrotendinous xanthometosis), polypeptide 1 (CYP27A1b) mRNA	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 65	Drosophila melanogaster merlin (Dmerlin) mRNA) complete cds	Chloken lipoprotein lipase gene	Chicken lipoprolein lipase gene	Grus canadensis recombination activating protein 1 (RAG-1) gene, partial cds	Grus canadensis recombination activating protein 1 (RAG-1) gene, partial cds	Bacillus halodurans genomic DNA, section 12/14	601067107F1 NIH_MGC_10 Hamo sapiens cDNA clone IMAGE:3453505 5	Cyanidium caldarium gene for SigC, complete cds	Oyanidium caldarium gene for SigC, complete cds
Top Hit Database Source	TN	M	TN	NT	SWISSPROT	NT	NT	Ŋ	N	EST_HUMAN		된	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	EST_HUMAN	<u>L</u>	F	N.	Į,	뒫	NT	NT	NT	EST_HUMAN	NT	ᅜ
Top Hit Acesslan No.	AF026198.1	X60986.1	AE003944.1	AE002186.2	026350	228337.1	D90911.1	AF106953.2	5901893 NT	AA595863.1		AF121970.1	AE004963.1	BF363970.1	BF107694.1	BF107694.1	AV661898.1	X17012.1	W69089.1	4503210 NT	AL161565.2	U48724.1	X60547.1	X60547.1	AF143732.1	AF143732.1	AP001518.1	BE542612.1	AB006799.1	AB006799.1
Most Similar (Top) Hit BLAST E Value	8.9E-01	8.9E-01			8.8E-01	8.8E-01	8.8E-01	8.7E-01	8.7E-01	8.7E-01		8.7E-01	8.7E-01	8.7E-01	8.7E-01	8.7E-01	8.7E-01	8.6E-01.	8.6E-01	8.65-01	8.6E-01	8.6E-01	8.6E-01	8.6E-01	8.6E-01	8.6E-01	8.6E-01	8.5E-01	8.5E-01	8.5E-01
Expression Signal	. 2.08	1.32	3.47	4.74	3.28	2.98	1.75	1.64	1.09	4.64		2.51	1.45	6.47	4.18	4.18	2.78	1.66	3.62	1.01	0.81	1.29	11.82	11.82	2.14	2.14	1.52	2.75	1.36	1.36
ORF SEQ ID NO:	25476		29085		24126					22560			27681			29088			20618	22012		23427	25586		26065					28014
Exan SEQ ID NO:	15413	15692				18304				12773		14816	17464	18090	18778	18776	19534	10409	10767	12109	13482	13642	15509		15934		16506	16304	17774	17774
Probe SEQ ID NO:	5494	5786	8990	9285	4442	8430	9108	457	2353	2845		4938	7613	8206	8970	8970	9488	466	840	2224	3568	3730	5855	5595	6030	6030	9299	6443	7924	7924

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סיישנים בייסים ב	Top Hit Descriptor	Homo sapiens human immunodeficiency virus type I enhancer-binding protein 1 (HIVEP1), mRNA	Rattus norvegicus protein tyrosine phosphatase, non-receptor type 5 (Ptpn5), mRNA	Mus musculus mper1 gene for period1, complete cds	Mus musculus NK cell receptor 2B4 gene, promoter region and partial cds	Human fibroblast growth factor receptor 3 (FGFR3) gene, Intron 7	Human fibroblast growth factor receptor 3 (FGFR3) gene, intron 7	Pyrococcus abyssi complete genome; segment 5/6	Thermus thermophilus cytochrome c-552 (cycA) and CycB (cycB) genes, complete cds	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 18	Nicotiana tabacum mRNA for chloroplast ribosomal protein L10, complete cds	Streptomyces antibioticus polyketide biosynthetic gene cluster	Canis familiaris MHC DLA Class II DRB pseudogene DRB2	Canis familiaris MHC DLA Class II DRB pseudogene DRB2	Canis familiaris MHC DLA Class II DRB pseudogene DRB2	Canis familiaris MHC DLA Class II DRB pseudogene DRB2	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 40	nn01f12.y5 NCI_CGAP_Co9 Homo saplens cDNA clone IMAGE:1076495 5' similar to contains THR.t1 THR	repetitive element ;	Drosophila melanogaster Lis1 homolog mRNA, complete cds	Mus musculus neuro-d4 gene, exons 3 through 12 and partial cds	Methanobacterium thermoautotrophicum from baises 1270510 to 1283409 (section 109 of 148) of the complete genome	Phytophthora infestans mitochandrion, complete genome	Homo sapiens FRA3B common fragile region. disidenosine triphosphate hydrolase (FHIT) gene, exon 5	Rettus norvegicus mRNA for RPHO-1, complete cds	Mus musculus trophinin (Tnn) gene, complete cds	IL3-CT0219-161199-031-C08 CT0219 Homo sapiens cDNA	Rattus norvegicus mRNA for RPHO-1, complete cds	CM4-HT0243-081199-037-e01 HT0243 Homo stiplens cDNA	S.cerevisiae MET, LEU4, and POL1 genes encoding MET4 protein, alpha-tsoproplymalate (alpha-IPM) synthetase (partial), and DNA polymerase alpha (partial)	Homo sapiens thioredoxin-related protein mRNA, complete cds	MCKUSICK-KAUFMAN/BARDET-BIEDL SYNDROMES PUTATIVE CHAPERONIN
IS LAUIT 1008	Top Hit Database Source			NT MU	NT	η. Hα			NT Th				NT Ca	NT C	N S	NT Ca	NT		T_HUMAN	NT	NT	Me		Ŷ.			T_HUMAN	NT	EST_HUMAN CN	S. TA		SWISSPROT MC
	Top Hit Acession No.	11418543 NT	9507008 NT	AB030818.1	AF143509.1	L78726.1	L78726.1	AJ248287.1	M93437.1	AL161506.2	AB010879.1		U46916.1	U46916.1	U46916.1	U46916.1	AL161540.2			AF098070.1	AF108133.1	A Enonger 4	2472	AF020503.1		Γ	AW376990.1	AB000489.1	AW379433.1	712128.1	1.6	09,1170
	Most Similar (Top) Hit BLAST E Value	8.5E-01	8.55-01	8.5E-01	8.4E-01	8.45-01	_				8.3E-01		8.3E-01	8.3E-01		8.3E-01	8.3E-01				8.3E-01	8 3E-04		8.3E-01	+	•				8.0F-01		8.2E-01
	Expression Signal	2.62	263	1.29	0.84	27	27	3.13	2.31	2.64	1	3.11	1.12	1.12	0.95	0.95	2.13		4.2	1.23	3.1	2.48	2.48	2.03	2.99	1.51	1.14	+	3.26	3.58	1.53	99.9
	ORF SEQ ID NO:			25243	23785	25145	25148			22774	23434	23619	24570	24571	24570	24571	24839			27914	27962	28204	10000	28760				24691		08890		28028
	Exon SEQ ID NO:	19643	19075	L	14008	19443		17571	10856			13839	14796	14796	14796	14796	15146		_		17717	17052	1	18488		L	12513	14917	16016	19462	ļ	17787
	Probe SEQ ID NO:	9430	9436	9683	4108	5378	5378	7721	724	3056	3739	3930	4917	4917	5177	5177	5222		7573	7822	7867	8082	8076	8623	2006	2043	2646	5045	6122	6300	7788	7837

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Top Hit Descriptor	MCKUSICK-KAUFMANIBARDET-BIEDL SYNDROMES PUTATIVE CHAPERONIN	Molluscum contaglosum virus type 1 ORF1 and ORF2 DNA	OVARIAN TUMOR LOCUS PROTEIN	yw14d02.r1 Soares_placenta_8to9weeks_2NbHF8to9W Homo sapiens cDNA clone IMAGE:252195.5" similar to ch:M38072 80S RIBOSOMAL PROTEIN L7A (HUMAN):	Mus musculus mRNA for NIPSNAP2 protein	Mus musculus TANK binding kinase TBK1 (Tbk1) mRNA, complete cds	Homo sapiens MHC class 1 region	Homo sapiens MHC class 1 region	NEURONAL MEMBRANE GLYCOPROTEIN M6-B	NEURONAL MEMBRANE GLYCOPROTEIN M6-B	RCO-TN0080-220800-025-d10 TN0080 Homo sepiens cDNA	RC0-TN0080-220800-025-d10 TN0080 Homo sepiens cDNA	Thermotoga maritima section 23 of 136 of the complete genome	Staphylococcus aureus partial pta gene for phosphate actyftransferase allele 15	Bos taurus futb and rtif genes	602072473F1 NCI_CGAP_Brn67 Homo sapiens cDNA clone IMAGE:4215091 5'	Salmiri boliviensis olfactory receptor (SBO27) gene, partial cds	Mus musculus gene for oviductal glycoprotein, complete cds	Nelsseria meningitidis serogroup A strain Z2491 complete genome; segment 7/7	G.gallus mRNA for nicotinic acetylcholine receptor (nAChR) beta 3 subunit	RC0-NN1012-270300-021-h06 NN1012 Homo saplens cDNA	Lymantria dispar nuclear polyhedrosis virus gene for DNA polymerase, complete cds	Ureaplasma urealyticum section 31 of 59 of the complete genome	Homo sapiens mRNA for KIAA1452 protein, partial cds	Haemophilus influenzae Rd section 54 of 163 of the complete genome	Oryctolagus cuniculus mRNA for mitsugumin29, complete cds	Danio rerio Trp4-associated protein Tap1A (tap1A) mRNA, complete cds	Gallus gallus SOX8 transcription factor (SOX8) niRNA, complete cds	601192033F1 NIH_MGC_7 Hamo sapiens cDNA clane IMAGE:3535785 5'	Mus musculus embigin (Emb), mRNA	Mus musculus embigin (Emb), mRNA	S.pneumoniae dexB, cap3A, cap3B and cap3C genes and orfs	S.pneumoniae ძომ, cap3A, cap3B and cap3C ganes and orfs
Top Hit Database Source	SWISSPROT	LN	SWISSPROT	NAMIH TRE	¥	Ę	Z	Z	SWISSPROT	SWISSPROT	EST_HUMAN	EST_HUMAN	Ę	Z	F	EST_HUMAN	TN	TN	NT.	ΙN	EST_HUMAN	TN	NT	TN	INT	TN	NT	F	EST_HUMAN	ΙΝ	۲	TN	N
Top Hit Acesslon No.	Q9JI70	L10127.1	P10383	H87308 1	AJ001261.1	AF191839.1	AF055066.1	AF055086.1	Q13491	Q13491	BE938558.1	BE938558.1	AE001711.1	AJ271510.1	AJ132772.1	BF530962.1	AF127897.1	AB006193.1	AL162758.2	X83739.2	AW901489.1	D11476.1	AE002130.1	AB040885.1	U32739.1	AB004816.1	AF130459.1	AF228684.1	BE263612.1	6753745 NT	6753745 NT	247210.1	247210.1
Most Similar (Top) Hit BLAST E Value	8.2E-01	8.2E-01	8.2E-01	8.2F_01	8.2E-01	8.1E-01	8.1E-01	8.1E-01	8.1E-01	8.1E-01	8.1E-01	8.1E-01	8.1E-01	8.0E-01	8.0E-01	8.0E-01	8.0E-01	8.0E-01	8.0E-01	8.0E-01	8.0E-01	7.9E-01	7.9E-01	7.9E-01	7.9E-01	7.9E-01	7.9E-01	7.9E-01	7.9E-01	7.9E-01	7.8E-01	7.9E-01	7.9E-01
Expression Signal	6.68	281	Θ	89.8	1.73	1.29	2.81	2.81	2.49	2.49	1.94	1.94	1.84	3.02	3.78	1.83	2.73	1.14	1.8	5.65	2.84	3.78	1.18	65.97	1.37	5.24	2.29	2.75	0.92	1.02	1.02	0.84	0.84
ORF SEQ ID NO:	Ц	29011		29069			23130		26020		28879		25343		20070		22761			24115		20211						23193		24188		24720	24721
Exon SEQ ID NO:		18717	18772	18777	L	12593	L	13329	15897		18592	18592	18909	10142	10250	11884		13194	13557	14327	16541		10631	11491	11538	12102		13388	14103	14401	14401	14945	14945
Probe SEQ ID NO:	7937	8809	9968	8971	9451	2731	3412	3412	2669	5992	8775	8775	9168	170	285	1990	3039	3273	3643	4432	6881	448	698	1587	1634	2216	2217	3472	4204	4508	4508	202	5075

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Single Exon Probes Expressed in Heart

	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLASTE Vatue	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
5084	14954		1.01	7.9E-01	7.9E-01 M29930.1	NT	Human insulin receptor (allele 2) gene, exons 14, 15, 16 and 17
6721	16601		2.43	7.9E-01	7.9E-01 X90996.1	NT	P.sativum GR gene
7504	1.7373	27582	4.78	7.9E-01	7.9E-01 U01912.1	TN	Glardia lamblia variant-specific surface protein G3M-B (vspG3M-B) mRNA, partial cds
7781	17631	27864	3.95	7.9E-01	P19719	SWISSPROT	SMALL HYDROPHOBIC PROTEIN
8372	18249		2.38	7.9E-01	7662471 NT	N.	Homo sapiens KIAA1072 protein (KIAA1072), mFiNA
8543		28682	2.48	7.9E-01	P19022	SWISSPROT	NEURAL-CADHERIN PRECURSOR (N-CADHERIN)
828	10784		1.75	7.8E-01	7.8E-01 Z43785.1	EST_HUMAN	HSC1KH041 normalized infant brain cDNA Homo sapiene cDNA clone c-1kh04
2229	12114			7.8E-01	97.1	EST_HUMAN	EST371637 MAGE resequences, MAGF Homo sapiens cDNA
4603	14491	24278	62.0	7.8E-01		NT	Rattus norvegicus transmembrane receptor Uncăl 1 mRNA, complete cds
5698	15607	25709	2:32	7.8E-01	7.8E-01 AF115856.1	벌	Sphenodon punctatus alpha enolase mRNA, partial cds
7323	17199	27399	1.27	7.8E-01	7.8E-01 Y10159.1	Ę	D.discoldeum racGAP gene
9424	19623		1.42	7.8E-01	7.8E-01 L29260.1	Ę	Arabidopsis thaliana 1-amino-1-cyclopropanecarboxylate synthase (ACS5) gene, complete cds
138	10112	19932	6.33	7.7E-01 AI	AF184345.1	ZT	Lycopersicon hirsutum ADP-glucose pyrophosphörylase large subunit (AGP-L1) mRNA, complete cds
							Mus musculus major histocompatibility locus class II règion: major histocompatibility protein class II alpha
							chain (IAalpha) and major histocompatibility protein class II beta chain (IEbeta) genes, complete cds;
709	10841		1.85		7.7E-01 AF050157.1	NT	butyrophilin-like (NG9), butyrophilin-li>
2680	12545	22438	2.8	7.7E-01	7.7E-01 033915	SWISSPROT	CITRATE SYNTHASE
0700	70007		•	20 75 5	114 007 0000	Ė	Homo sapiens UDP-N-acetyk-alpha-D-galactosamine:polypeptide N-acetylgalactosaminytransferase 7
2 20	1				A 174 4000	F 12	(Camaron I) (Control I) (Camaron I) (Camar
Race				A10-27.7	AF 1 10000.1	<u> </u>	rigino separa e e e e e e e e e e e e e e e e e e
4297	14195			7.7E-01	7.7E-01 AF199488.1	Z.	Cournix cournix japonica sub-species japonica beta-actin mixiva, partial cos
4297	L	23980		7.7E-01	7.7E-01 AF199488.1	LN	Courink courink aponica sub-species Japonica bera-actin mkNA, partial cos
5415				7.7E-01	P16553	SWISSPROT	RAFFINOSE INVERTASE (INVERTASE)
5415				7.7E-01	P16553	SWISSPROT	RAFFINOSE INVERTASE (INVERTASE)
5628	15541	25630	1,95	7.7E-01	7.7E-01 R08600.1	EST_HUMAN	yr24b02.s1 Soares felal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:1277553'
9212	10112	19932	2.8	·	7.7E-01 AF184345.1	Z	Lycopersican hirsutum ADP-glucose pyrophosphorylase large subunit (AGP-L1) mRNA, complete cds
9311	1				11497621 NT	۲	Archaeoglobus fulgidus, complete genome
							Arabidopsis thallana 3-methylcrotonyl-CoA carboxylase non-biotinylated subunit (MCCB) mRNA, complete
5710	15618	25720	4.04	7.6E-01 A	AF059510.1	Ę	sþo
6710	15618	25721	4:04	7.8E-01	AF059510.1	LN	Arabidopsis thaliana 3-methylcrotony/-CoA carboxylase non-biotinylated subunit (MCCB) mRNA, complete cds
	1						

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Probe SEQ ID NO:	Exen SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acesslan No.	Top Hit Database Source	Тф Hit Descriptor
6699	16579	26770	1.41	7.6E-01	AF146793.2	F	Mus musculus neuromedin U precursor (Nmu) gene, partial cds; tPhLP (Tphip) gene, partial cds; CLOCK (Clock) gene, complete cds; PFT27 (Pf27) gene, complete cds; and H5AR (H5ar) gene, complete cds
6726	16806	26796	2.04	7.6E-01	6867762 NT	ᅜ	Mus musculus advillin (Advil-pending), mRNA
6728	16606	26797	2.04	7.6E-01	6857752 NT	뒫	Mus musculus advillin (Advil-pending), mRNA
7184	17061	27252	1.28		6753577 NT	Ы	Mus musculus cytochrome P450, 2b9, phenobarbitol Inducible, type a (Cyp2b9), mRNA
7347	17215	27414	7.52	,	P30372	SWISSPROT	MUSCARINIC ACETYLCHOLINE RECEPTOR M2
7347	17215	27415		7.6E-01	7.6E-01 P30372	SWISSPROT	MUSCARINIC ACETYLCHOLINE RECEPTOR M2
8664	18553	28837	2.31	7.6E-01	X86347.1	L.	H.aspersa mRNA for neurofilament NF70
8664	18553	28838	2.31	7.6E-01	X86347.1	NT	H.aspersa mRNA for neurofilament NF70
8948	18756		4.81	7.6E-01	7.6E-01 AL161592.2	TN	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 88
9073	18850		2.74		AB020702.1	TN	Homo sapiens mRNA for KIAA0895 protein, partial cds
602	10444		1.62		7.5E-01 AL163301.2	NT	Homo sapiens chromosome 21 segment HS21C101
999	10508	20315	2.23	7.5E-01	7.5E-01 AF020503.1	Į.	Homo sapiens FRA3B common fragile region, diadenosine triphosphate hydrolase (FHIT) gene, exon 5
3315	ı				C14203.1	EST_HUMAN	C14203 Clontech human aorta polyA+ mRNA (#6572) Homo saplens cDNA clone GEN-037E11 5'
4573	10083	18900	13.92		L	M	Homo sapiens hypothetical protein FLJ10793 (FLJ10793), mRNA
9381	19042		3.57	7.5E-01	AF163151.2	NT	Homo sapiens dentin statophosphoprotein precursor (DSPP) gene, complete cds
9828	19329	25209	1.87	7.5E-01	D90907.1	TN	Synechocystis sp. PCC6803 complete gename, 9/27, 1056467-1188885
9884	19368	25190	1.42	7.5F-01 A	AE000823.1	Þ	Methanobacterium thermoautotrophicum from basis 317350 to 328792 (section 29 of 148) of the complete denome
							h14b09.x1 NCI_CGAP_Bm25 Homo saplens cDNA clone IMAGE:2167577 3' similar to contains Alu
1114	11029		1.23	7.4E-01	7.4E-01 AI598146.1	EST_HUMAN	repetitive element contains element MIR repetitive element;
2295	12177	22076	0.97	7.4E-01	7.4E-01 AB011106.1	IN	Homo sapiens mRNA for KIAA0534 protein, partial cds
3665	13579	23366	1.22	A 10-34.7	AF112538.1	NT	Malva pusilla actin (Act1) mRNA, complete cds
4213	14111	23889	4.43	7.4E-01	AL163246.2	ΙN	Homo sapiens chromosome 21 segment HS21C048
7258	17135	27328	7.23	7.4E-01	BE74750	EST_HUMAN	601573028F1 NIH_MGC_9 Hamo sapiens cDNA clone IMAGE:3834174 5'
9041	18828		3.28		6753217 NT		Mus musculus complement component 1 inhibitor (C1nh), mRNA
9150	18901		1.59		7.4E-01 AI472841.1	EST_HUMAN	ta13h01.x1 NCI_CGAP_Lym5 Homo saplens cDNA clone IMAGE:2043985 3'
4515	14408	24194	96.0	7.3E-01	AE001166.1	MT	Borrelia burgdorferi (section 52 of 70) of the complete genome
4602	14490	24277	4.78	A 10-3E-01	AF225421.1	NT	Homo sapiens HT017 mRNA, complete cds
5127	14994		1.1	7.3E-01	1 1	ᅜ	Homo sapiens growth arrest-specific 7 (GAS7), transcript variant b, mRNA
5984		26010				뉟	Mus musculus antigen (CD72) gene
5984	15889		5.61	7.3E-01	135772.1	본	Mus musculus antigen (CD72) gene

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Probe SEQ ID

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6454 6454 8729 8729 1914 2412

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Homo saplens transortption factor IGHIM enhancer 3, JM11 protein, JM4 protein, JM5 protein, T54 protein, JM10 protein, A4 differentiation-dependent protein, triple LIM domain protein 6, and synaptophysin genes, Homo sapiens transcription factor IGHIM enhancer 3, JM11 protein, JM4 protein, JM5 protein, T54 protein, JM10 protein, A4 differentiation-dependent protein, thoie LIM domain protein 6, and synaptophysin genes, Rana catesbeiana mRNA for bullfrog skeletal muscle calcium release channel (ryanodine receptor) alpha Giardia intestinalis variant specific surface protein (vsp417-6) gene, vsp417-8/A-I allele, complete cds 2/25b08.s1 Soares_fetal_liver_spleen_1NFLS_S1 Homo saplens cDNA clone IMAGE:431799 3' 2/25b08.s1 Soares_fetal_liver_spleen_1NFLS_S1 Homo saplens cDNA clone IMAGE:431799 3' Drosophila melanogastar 6-pyruvoyltetrahydropterin synthase (pr) gene, complete cds 801498330F1 NIH_MGC_70 Homo saplens cDNA clone IMAGE;3898495 5 602035589F1 NCI_CGAP_Bm64 Homo sapiens|cDNA clone IMAGE:4183222 5 Homo saplens partlal TCF-4 gene for T-cell transcription factor-4, exons 15-16 zu06h11.s1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:731109 3 Oryctolagus cuniculus RING-finger binding protein mRNA, partial cds 602118381F1 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:4275381 5 602155438F1 NIH_MGC_83 Homo sapiens cDNA done IMAGE:4296344 5' 602155438F1 NIH_MGC_83 Homo sapiens cDNA clone IMAGE:4296344 5' Rattus norvegicus initiation factor-2 kinase (eIF-2a) mRNA, complete cds Top Hit Descriptor L.mesenteroides gene for sucrose phosphorylase (EC 2.4.1.7) Gallus gallus gene for melanocortin 2-receptor, complete cds Homo saplens mRNA for KIAA0614 protein, partial ods Homo sapiens mRNA for KIAA0614 protein, partial cds Rettus norvegicus cytocentrin mRNA, complete cds Aeropyrum pernix genomic DNA, section 6/7 V. alginolyticus sucrase (scrB) gene, complete cds V.alginolyticus sucrase (scrB) gene, complete cds complete cds; and L-type calcium channel a> complete cds; and L-type calcium channel a> Mus musculus otogelin (Otog), mRNA Mus musculus otogelin (Otog), mRNA Fowlpox virus, complete genome N.tabacum NelF-4A13 mRNA soform(RyR1), complete cds **EST HUMAN** EST_HUMAN HUMAN EST_HUMAN EST HUMAN **EST HUMAN** HUMAN EST_HUMAN Top Hit Database Source EST EST 눋 Ę 뉟뉟 뉟 눋 눋 눋 눋 둗 눋 F 눋 눋 Top Hit Acessian 7.1E-01 BF681034.1 7.1E-01 AA421492.1 7.0E-01 AB014514.1 7.2E-01 AF196779.1 7.2E-01 BF670061.1 7.3E-01 AA678019.1 .3E-01 AA678019.1 7.2E-01 AF065606.1 BF338350.1 7.2E-01 AF196779.1 7.2E-01 AF198100. 7.1E-01 AJ270777.1 BF681034.1 BE904405.1 7.2E-01 AB009605. AP000063. ģ 7.1E-01 U36232.1 7.2E-01 D90314.1 7.2E-01 129281.1 .2E-01 U82623.1 X79140.1 **D21070**. 2E-01 7.2E-01 7.3E-01 7.1E-01 7.1E-01 7.1E-01 15-01 .1E-01 (Top) Hit BLAST E Value 3.29 3.29 25. 2.14 1.16 10.38 1.48 108 2.93 1.56 1.04 1.04 3.68 2.41 222 4.08 1.58 6.97 1.61 Expression Signal 26481 28870 23794 20966 23124 23494 24707 26994 25625 22186 22748 24335 24706 26162 ORF SEQ 28871 21687 28258 ö 2 SEQ ID 16315 11118 18316 13323 14934 10610 19550 18585 18685 11809 12289 12956 14546 14934 18009 19160 12952 15538 16022 -10742 14014 14014 13707 16801 ÿ

5064

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			_	т-	_			Τ.	- -	_			_	_	_	_	_	_	_		_	-		_	_	_	_	
Top Hit Descriptor	yz73e07.s1 Soares_multiple_sclerosis_ZNbHMSP Homo sapiens cDNA clone IMAGE:288708 3' similar to contains Alu repetitive element;	yz73e07.s1 Soares_multiple_sclerosis_ZNbHM3P Homo sapiens cDNA clone IMAGE:288708 3' similar to contains Alu repetitive element;	Homo sapiens chromosome 21 segment HS21C101	Escherichia coli K-12 MG1655 section 143 of 400 of the complete genome	AV763842 MDS Homo sepiens cDNA clone MDSCHE04 5'	AV763842 MDS Homo sapiens cDNA clone MDSCHE04 5'	Candida albicans squalene epoxidase (CAERG1) gene, complete cds and translational regulator gene, partial cds	Candida albicans squalene epoxidase (CAERG1) gene, complete cds and translational regulator gene, partial	cds	nn28a09.s1 NCI_CGAP_Gas1 Homo saplens ciDNA clone IMAGE:1085176 3'	Chlamydia muridarum, section 3 of 85 of the complete genome	Mus musculus mRNA for immunoglobulin gamma heavy chain variable region, isolate PC 2811	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 69	Arabidopsis thaliana DNA chromosome 4, contigifragment No. 69	Homo sapiens DAN gene, complete cds	Homo sapiens DAN gene, complete cds	FORKHEAD BOX PROTEIN C2 (FORKHEAD-FIELATED PROTEIN FKHL14) (MESENCHYME FORK HEAD PROTEIN 1) (MFH-1 PROTEIN) (TRANSCRIPTION FACTOR FKH-14)	Glardia intestinalis carbamate kinase gene, complete cds	Synechocystis sp. PCC6803 complete genome, 27/27, 3418852-3573470	aj75a05.s1 Soarss_parathyroid_tumor_NbHPA Homo sapiens cDNA clone IMAGE:1402256 3' similar to gb:X56411_rna1 ALCOHOL DEHYDROGENAS;; CLASS II PI CHAIN (HUMAN);	Rat(hooded) prolactin gene : exon ill and flanks	Homo sapiens hevin (HEVIN) mRNA	Homo sapiens mRNA for KIAA1345 protein, partitul cds	Stagonospora avenae bgl1 gene for beta-glucosidase, exons 1-4	Stagonospora avenae bgl1 gene for beta-glucosidase, exons 1-4	Mus musculus zinc finger protein (Peg3) mRNA, complete cds	Mus musculus zinc finger protein (Peg3) mRNA, complete cds	Anopheles gambiae strain M2 translation Initiation factor 4C (1A) (eIF-4C) mRNA, complete cds
Top Hit Database Source	EST_HUMAN	EST HUMAN	N L	LZ.	EST_HUMAN	EST_HUMAN	Ę		٦	EST_HUMAN	TN	TN	F	۲	TN	IN	SWISSPROT	Ę	Z	EST_HUMAN	Ā	LΝ	뉟	Z	LN	NT	NT	N
Top Hit Acession No.	N62412.1	N62412.1	AL163301.2	AE000253.1	AV763842.1	AV763842.1	U69674.1		U69674.1	AA593530.1	AE002271.2	Y17373.1	AL161573.2	AL161573.2	D89013.1	D89013.1	Q99958	AF017784.1	D90917.1	AA854475.1	J00762.1	4758521 NT	AB037766.1	AJZ76675.1	AJ276675.1	AF038939.1	AF038939.1	AF164151.1
Most Similar (Top) Hit BLAST E Value	7.0E-01	7.0E-01	7.0E-01	7.0E-01	7.0E-01	7.0E-01	6.9E-01		6.9E-01	6.9E-01	6.9E-01	6.9E-01	6.9E-01	6.9E-01	6.9E-01	6.9E-01	6.9E-01	6.8E-01	6.8E-01	6.8E-01	6.8E-01	6.8E-01	6.8E-01	6.8€-01	6.8E-01	6.8E-01	6.8E-01	6.8E-01
Expression Signal	4.	4.1	1.95	8.1	2	2	15.73		15.73	2.21	1.63	0.85	2.73	2.73	3.58	3.56	237	1.03	1.16	1.58	1.26	0.83	1.52	2.34	234	2.59	2.59	1.79
ORF SEQ ID NO:	22173	22174				28600	20725					23117	26728	28727	28719	28720		20713		21362	24153	24420	27622	28566	28567	28593		
Exon SEQ ID NO:	1227	12277		16759	18336	18336	10878		10878	11186		13316	16532	16532	18450	18450	19543	10866	12508	11502		14634	17407	18311	18311	18331		18485
Probe SEO ID NO:	2399	2399	4989	6880	8463	8463	954		954	1288	3182	3399	6652	6652	8682	8582	9016	941	2639	2799	4469	4749	7556	8437	8437	8428	8458	8619

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Table 4
Single Exon Probes Expressed in Heart

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יינוסטיין ויינוסטיין דיין ויינוסטיין דיין ויינוסטיין דיין ויינוסטיין דיין ויינוסטיין דיין ויינוסטיין דיין ויינוסטיין	Top Hit Descriptor	Homo sapiens nuclear factor of kappa light polypeptide gene enhancer in B-celle 1 (NFKB1) gene, complete ods	Homo sapiens nuclear factor of kappa light polypeptide gene enhancer in B-cells 1 (NFKB1) gene, complete ods	ex12g12.s1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA clone IMAGE:786310 3' similar to contains element TAR1 repetitive element :	Drosophila melanogaster Mst85C gene, complete cds; NMDMC Isoform (Nmdmo) gene, complete cds, alternatively spliced; and transoription factor (Relish) gene, complete cds, alternatively spliced	Mus musculus Wiskott-Aldrich syndrome protein (Wasp), mRNA	S.tuberosum mRNA for glucose-6-phosphate dehydrogenase	xe95g12.x1 NCI_CGAP_Co17 Homo septens cDNA clone IMAGE:2574598 3'	Dendrobium fimbriatum mRNA for phosphoenolpyruvate carboxylase, partial	Helicobacter pylori, strain J99 section 47 of 132 of the complete genome	Gallid herpesvirus 2, complete gename	Gallid herpesvirus 2, complete genome	Pseudomonas aeruginosa PA01, section 167 of 529 of the complete genome	Helicobacter pylori, strain J99 section 47 of 132 of the complete genome	CM3-HT0769-010600-197-c03 HT0769 Homo saplens cDNA	N-ACETYLGLUCOSAMINYL-PHOSPHATIDYLINOSITOL BIOSYNTHETIC PROTEIN GPI1	Homo sapiens SLIT1 protein (SLIL2) mRNA, partial cds	Homo sapiens lens epithelium-derived growth factor gene, alternatively spliced, complete cds	Homo saplens sema domain, seven thrombosporidin repeats (type 1 and type 1-like), transmembrane domain (TM) and short cytoplasmic domain, (semaphorir) 5A (SEMA5A) mRNA	Cabicans random DNA marker, 282bp	Human hereditary haemochromatosis region, histone 2A-like protein gene, hereditary haemochromatosis (HLA-H) gene, RoRet gene, and sodium phosphete transporter (NPT3) gene, complete cds	Mus musculus kinesin light chain 2 (Kic2), mRNA	AV660506 GLC Homo sapiens cDNA clone GLCGID04 3'	Homo sapiens chromosome 21 segment HS21C078	Vibrio cholerae chromosome II, section 39 of 93 of the complete chromosome	H.vulgaris Na,K-ATPase alpha subunit mRNA, complete cds	H.vulgaris Na.K.ATPase alpha subunit mRNA, complete cds
01 1 100-1016	. Top Hit Database Source	Ę	뒫	EST_HUMAN	IN	LΝ	TN	EST_HUMAN	NT	NT	L	, LN	NT	. LN	EST_HUMAN	SWISSPROT	IN	IN	LΝ	TN	· L	F	EST HUMAN	F	NT	NT	L
5	Top Hit Acession No.	AF213884.1	AF213884.1	AA451864.1	AF186073.1	FR 6678580 NT	X74421.1	AW079110.1	AJ252842.1	AE001486.1	9635035 NT	9635035 NT	AE004606.1	AE001486.1	BF354649.1	014357	AF075240.1	AF199339.1	4608880 NT	Y07669.1	U91328.1	6680577	AV660508.1	AL163278.2	AE004382.1	M75140.1	M75140.1
	Most Similar (Top) Hit BLAST E Value	6.7E-01	8.7E-01	8.7E-01	6.7E-01	6.7E-01		6.7E-01			6.7E-01	6.7E-01	6.7E-01		6.7E-01	6.7E-01		6.6E-01	6.6E-01	6.6E-01	8.6E-01	_			6.6E-01		6.5E-01
	Expression Signal	29.34	19.53	1.57	3.59	3.7	96.0	0.85	0.89	1.62	1.42	1.42	3.98	1.46	2.23	3.69	0.87	1.29	1.04	3.05	0.87	3.86	3.63	2.41	1.35	1.37	1.37
	ORF SEQ ID NO:	20080	20110	21887	21908	22692	24032	24538			25838	25837		26367	28449			22422	23168			25840			25256		20353
	Exan SEQ ID NO:	10259	10295	11989	12713	12893	14246	14762	15048		15723	15723	16192	1	18200	17803		12532	13362	13522	13924			1			10544
	Probe SEQ ID NO:	295	338	2100	2120	2966	4350	4881	5182	5627	5817	5817	6329	6342	8323	8754	2452	2867	3445	3608	4020	5822	6525	7571	9619	809	809

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Table 4
Single Exon Probes Expressed in Heart

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	Exon SEQ ID NO:	ORF U	Expression Signal	Most Similar (Top) Hit BLAST E Value	,	Top Hit Database Source	Top Hit Descriptor
3386	13304	23104	6.1	6.5E-01	AB041225.1	NT	Mus musculus gene for Tob2, complete cds
3951	13859	23633	1.08	6.5E-01	4504632 NT	NT	Homo sapiens interleukin 10 receptor, alpha (III10RA) mRNA
4185	14085		4.17		AJ272265.1	뒫	Homo sapians SPP2 gene for secreted phosphoprotein 24 precursor, exons 1-8
. 4215	14113	23890	0.86		AL161539.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 39
4994	14869	24632	2.27	6.5E-01	U28921.1	ŢN	Phaseolus vulgaris ATPase gamme subunit mRNA, nuclear gene encoding mitochondrial protein, partial cds
A178	15041		88.0		1137258 1	Ę	Acetobacter xylinum putative ATP binding protein delta-AceB gene, partial cds, and GDP-mannose:cellobiosyldinhosobocoloxieno alobe-mannosyltransferse complete cds.
96036	15939	26071	1.38			Z	Ohicken mRNA for 115-kDa melanosomal matrix protein, complete cds
7919	17769			6.5E-01	AF119676.1	NT	Mus musculus small GTP-binding protein RAB25 (Rab25) gene, complete cds
0608	17922		3.24			EST_HUMAN	yw17106.r1 Soares_placenta_8tc9weeks_2NbH;>8tc9W Homo sapiens cDNA clone IMAGE:252515 5'
8073	17864			6.5E-01	AA601287.1	EST_HUMAN	no15c07,s1 NCI_CGAP_Phe1 Homo saplens cDNA clone IMAGE:1100748 3'
8170	18058		4.27	L	AU138078.1	EST_HUMAN	AU138078 PLACE1 Homo sapiens cDNA clone PLACE1007810 5'
8877	18689	28981	23	6.5E-01	AF014115.1	TN	Plasmodium berghei cytochrome c oxidase subt ⁱ rit III, cytochrome c oxidase subunit I, and cytochrome b genes, mitochondrial genes encoding mitochondrial proteins, complete cds
9419	19069		2.79	8.5E-01	BE465050.1	EST_HUMAN	hv74a10.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3179130 3'
9851	19490		1.69	6.5E-01	Z74145.1	NT	S.cerevisiae chromosome IV reading frame ORF YDL097c
252	10218	20035	5.4	6.4E-01	U48848.1	NT	Drosophila melanogaster 8kd dynein Ilght chain inRNA, complete cds
3410	13327	23128	2.45	6.4E-01	ĭ	NT	Mus musculus dystroglycan 1 (DAG1) gene, excirs 1 and 2 and complete cds
3785	13697	23484	1.34	6.4E-01	AB046827.1	NT	Homo saplens mRNA for KIAA1607 protein, partial cds
4389	14285		0.84		Y12488.1	NT	M.musculus whn gene
4389	14285	24067	0.84	6.4E-01	Y12488.1	INT	M.musculus whn gene
. 7007	16884	27076	1.76		1	TN	Treponema pallidum section 63 of 87 of the complete genome
7807	17657	27895	13.62	6.4E-01	1	TN	Homo sapiens ataxla telangiectasia (ATM) gene, complete cds
7818	17668	27908	1.44	6.4E-01	1	EST_HUMAN	602150289F1 NIH_MGC_81 Homo sapiens cDNA clone IMAGE:4291126 5'
9528	19135		6.34	. 6.4E-01	AV75921	EST_HUMAN	AV759212 MDS Homo sapiens cDNA clone MDSCGC09 5
9951	19553		1.65	6.4E-01	9845300 NT	NT	Rat cytomegalovirus Maastricht, complete genome
427	10372	20198	4.41		_	SWISSPROT	HISTIDINE-RICH PROTEIN PRECURSOR (CLONE PFHRP-III)
524	10466	20277	2.15	6.3E-01	U32689.1	TN	Haemophilus influenzae Rd section 4 of 163 of the complete genome
2116	12005		3.87	6.3E-01		NT	Shigella flexneri multi-antiblotic resistance locus
2534	12408	22300				INT	Gallus gallus bone morphogenetic protein 1 (BMP ₁ 1) mRNA, partial cds
2534	12408			6.3E-01		뒫	Gallus gallus bone morphogenetic protein 1 (BMP ₁ 1) mRNA, partial cds
2883	12911		0.83		Y17275.1	Į.	Lycopersican esculentum p69a gene, complete CDS

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Probe SEQ ID S NO:	Exen SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Tcp Hit Descriptor
3985	13892	23668	1.38	6.3E-01	X99675.1	NT	D.melanogaster mRNA for metabotropic glutamate receptor
6369	16837		3.17	6.3E-01	BE902044.1	EST_HUMAN	601876889F1 NIH_MGC_21 Homo saplens cDŅA clone IMAGE:3959351 5'
7150	17027	27223	1.67	6.3E-01	S62927.1	IN	glycoprotein IIIa (Alu 1 and 3 fusion junction) [human, Genomic Mutant, 300 nt]
7421	17288	27495	2.72	6.3E-01	9627521 NT	NT	Variola virus, complete genome
7421	17288	27496	2.72	6.3E-01	1257529 NT	NT .	Variola virus, complete genome
8004	17854	28095	1.4	6.3E-01	AE000313.1	NT	Escherichia coli K-12 MG1655 section 203 of 400 of the complete genome
8443	18288	28543	70.0	A 3F.01	0 A B 7 7 7 7 4 4	NAMILH TRA	nn09h06.s1 NCI_CGAP_Co10 Home sepiens cDNA clone IMAGE:1181371 3' similer to TR:002916 002916 HI ARK
8647	18511	28791	11.72	6.35-01		EST HUMAN	CM-BT043-090299-046 BT043 Homo sapiens clDNA
8725	18581	28865	1.94	6.3E-01		SWISSPROT	HYPOTHETICAL 13.7 KD PROTEIN IN INO1-IDS2 INTERGENIC REGION
8888	18680	28969	1.98	6.3E-01	P36073	SWISSPROT	HYPOTHETICAL 15.3 KD PROTEIN IN VMA12-APN1 INTERGENIC REGION
9128	19697	24899	9.21	8.3E-01	PN 5820166	Ę	Mus musculus keratin complex 2, gane 6g (Krt2-6g), mRNA
9220	18944		1.54	6.3E-01	AF105227.1	NT	Homo sapiens 3'-phosphoadenosine 5'-phosphosulfate synthetase (PAPSS) mRNA, complete cds
9435	19819		1.58	6.3E-01	X83528.1	M	C.limicola pscD gene
2278	15493	25569	2.12	6.2€-01	Q10135	SWISSPROT	HYPOTHETICAL 142.5 KD PROTEIN C23E2.02 IN CHROMOSOME I
6426	16287		3.06	6.25-01		TN	Mus musculus calcium-sensing receptor related protein 4 (Casr-rs4) mRNA, partial cds
6831	18710	26903	5.45	6.2E-01	H72255.1	EST_HUMAN	ys01e08.s1 Soares fetal liver spleen 1NFLS Homo saptens cDNA clone IMAGE:213542 3'
7435	16448	26638	1.71	6.2E-01		EST_HUMAN	601336146F1 NIH_MGC_44 Homo saplens cDNA clone IMAGE:3690010 5
7476	17336		2.56	6.2E-01	M24461.1	NT	Human pulmonary surfactant-associated protein SP-B (SFTP3) mRNA, complete cds
7799	17649	27886	7.14	6.2E-01	AL161511.2	NT	Arabidopsis thallana DNA chromosome 4, contig fragment No. 23
8007	17857	28099	5.02	6.25-01	P27410	SWISSPROT	NON-STRUCTURAL POLYPROTEIN [CONTAINS: RNA-DIRECTED RNA POLYMERASE ; THIOL PROTEASE P3C ; HELICASE (2C LIKE PROTEIN); COAT PROTEIN]
8007	17857	28100	5.02	6.25-01	P27410	SWISSPROT	NON-STRUCTURAL POLYPROTEIN (CONTAINS: RNA-DIRECTED RNA POLYMERASE; THIOL PROTEASE P3C: HELICASE (2C LIKE PROTEIN); COAT PROTEIN)
2345	12225		4.5	6.1E-01		Į.	Mus muscullus secreted acidic cysteine rich glycoprotein (Sparc), mRNA
4931	14809	24577	0.99	6.1E-01	L20427.1	Ę	Rattus norvegicus dihydroxypolyprenylbenzoate methyltransferase mRNA, complete cds
4931	14809	24578	66.0	6.1E-01	1.20427.1	F	Rattus norvegicus dihydroxypolyprenylbenzoate methyltransferase mRNA, complete cds
. 6104	15998	26134	3.78	6.1E-01		NT	Rat TRPM-2 gene, complete cds
6104	15998	26136	3.78	6.1E-01	M64733.1	NT	Rat TRPM-2 gene, complete cds
6792	16871	28863	3.72	6.1E-01	AF0335	NT	Arabidopsis thaliana putative zinc transporter (ZIP1) mRNA, complete cds
	16985		1.75	6.1E-01	11431065 NT	٦	Homo sapiens mitogen-activated protein kinase kinase kinase 4 (MAP4K4), mRNA
il	16985		1.75	. 6.1E-01	11065	Į.	Homo sapiens milogen-activated protein kinase kinase kinase 4 (MAP4K4), mRNA
7419	17286	27492	19.47	8.1E-01		ᅜ	Homo sepiens G-protein coupled receptor EDG-7 mRNA, complete cds
7419	17286	27483	19.47	6.1E-01	AF236117.1	ĮŅ.	Homo saplens G-protein coupled receptor EDG-7 mRNA, complete cds

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	Top Hit Descriptor	Homo sapiens dopamine transporter (SLC6A3) gene, complete cds	Homo sapiens DNA for amyloid precursor protein, complete cds	Homo sapiens adaptor-related protein complex 3, mu 2 subunit (CLA20), mRNA	Human respiratory syncytial virus strain CH93-53b attachment protein (G) gene, complete cds	Viral hemorrhagic septicemia virus N, P, M, G, Nv, L genes, French strain 07-71	D(2) DOPAMINE RECEPTOR	UI-H-BI1-aab-a-10-0-UI.s1 NCI_CGAP_Sub3 Homo sapiens cDNA clone IMAGE:2718619 3'	Musca domestica insecticide-susceptible strain voltage-sensitive sodium channel mRNA, complete cds	Homo sapiens partial LMO1 gene for LIM domain only 1 protein, exon 1	SEGMENTATION PROTEIN FUSHI TARAZU	SEGMENTATION PROTEIN FUSHI TARAZU	ane B4 receptor BLT1, complete cds	PEROXISOMAL MEMBRANE PROTEIN PERS((PEROXIN-3)	Gallus gallus mRNA for Hyperion protein, 419 kD isoform	Gallus gallus mRNA for Hyperion protein, 419 kD isoform	#108f07.x1 NOL_CGAP_Pr28 Homo septens cDNA clone IMAGE:2095621 3'	Homo sapiens nuclear factor (erythroid-derived 2)-like 3 (NFE2L3), mRNA	Mus musculus cGMP-inhibited phosphodlesterase (Pde3a), mRNA	RC1-HT0375-030500-015-c03 HT0375 Homo sepiens cDNA	Haemophilus influenzae Rd section 16 of 163 of the complete genome	Homo sapiens chromosome 21 segment HS21C067	Homo sapiens chromosome 21 segment HS21C067	Rattus norvegicus cenexin 2 mRNA, partial cds	Homo sepiens low density lipoprotein receptor-related protein II (LRP2) gene, exon 1 and partial cds	THYMIDYLATE KINASE (DTMP KINASE)	PM1-DT0041-190100-002-h03 DT0041 Homo sapiens cDNA	Mus spretus strain SPRET/El CD48 antigen (Cd48) gene, partial cds	Oryctolagus cuniculus alpha 1 anti-trypsin (alpha 1 AT) gene, promoter region	Aspergillus oryzae pyrG gene for orolidine-5-phosphate decarboxylase, complete cds	MICROTUBULE-ASSOCIATED PROTEIN 1A [CONTAINS: MAP1 LIGHT CHAIN LC2]	SIM1 PROTEIN
	Top Hit Database Source	NT	NT	· LN	IN	TN	SWISSPROT	EST_HUMAN	NT	ΤŅ	SWISSPROT	SWISSPROT	LΝ	SWISSPROT	TN	NT	EST_HUMAN	M	Ā	EST_HUMAN	M	뒫	뒫	Ψ	F	SWISSPROT	EST_HUMAN	卢	ᅜ	TN.	SWISSPROT	SWISSPROT
	Top Hit Acession No.	AF119117.1	D87675.1	5802999 NT	AF065253.1	AJ233396.1	P20288	AW139713.1	U38813.1	AJ277661.1	P02835	P02835	AB008193.1	Q01497	AJ131892.1	AJ131892.1	A1420623.1	11421663 NT	9055303 NT	BE157617.1	U32701.1	AL163267.2	AL163287.2	AF162758.1	AF065440.2	Q9X0[3	AW937175.1	AF064626.1	L42320.1	AB017705.1	P34926	P40472
	Most Similar (Top) Hit BLAST E Value	6.1E-01	6.0E-01	6.0E-01	6.0E-01	6.0E-01	6.0E-01	6.0E-01	6.0E-01	6.0E-01	6.0E-01	6.0E-01	6.0E-01	6.0E-01	6.0E-01	6.0E-01	6.0E-01	6.0E-01	6.0E-01					5.9E-01	5.9E-01	5.9E-01	5.9E-01		6.9E-01		6.9E-01	5.8E-01
	Expression Signal	1.59	0.92	2.64	1.78	1.02	1.58	2.58	2.61	6.2	4.19	4.19	1.69	1.43	1.78	1.78	2.74	1.84	2.1	2.18	1.25	4.85	4.85	4.12	2.1	2.67	3.1	2.36	1.91	2.18	4.15	1.44
	ORF SEQ ID NO:	27860	20243		21103		24918	25086	25968	26375	26794	26795	27727		28541	28542		25291	25065		20752				25917	28203	28454		25342			21637
	SEQ ID NO:	17628	10429	10490	11245		15151	15260	15845	16213	16605	16605	17503	17744	18286	18286		19115	19554	19484	Ш		13156	14029	15795	17952	18205				19183	11763
	Probe SEQ ID NO:	7778	486	548	1339	3744	5227	5339	5940	6350	6725	6725	7653	7894	8411	8411	8838	9500	9787	6086	984	3232	3232	4129	5889	8061	8328	8529	9165	9404	9614	1867

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Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Vatue	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
3902	13812	23596	96'0		BF695738.1	EST_HUMAN	601852474F1 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:4076131 5
4413	14307		2.81	5.8E-01	AB009077.1	NT	Vigna radiata mRNA for proton pyrophosphatase, complete cds
4690	14576		1.04	5.8E-01	AF110846.1	NT	Megaselia scalaris sex-lethal homolog (MegsxI) gene, partial cds, alternatively spliced products
5753	15661	25768	1.29		D78659.1	EST_HUMAN	HUM500E06B Human placenta polyA+ (TFujiwara) Homo sapiens cDNA clone GEN-500E06 5
6077	16060		2.3	5.8E-01	S65091.1	NT	cyclic AMP-regulated phosphoprotein [rats, mRNA, 1030 nt]
6804	16484		2.62	5.8E-01	H41571.1	EST HUMAN	yn91b03.s1 Soares adult brain N2b5HB55Y Homo sapiens cDNA clone IMAGE:175757 3' similar to ob:S78187 M-PHASE INDUCER PHOSPHATASE 2 (HUMAN):
6764	16843	26830	2.23	5.8E-01		SWISSPROT	SPORE COAT PROTEIN SP96
6764	16643	26831	2.23		P14328	SWISSPROT	SPORE COAT PROTEIN SP98
7153	17030		8.64	5.8E-01	AJ270774.1	NT	Homo sapiens partial TCF-4 gene for T-cell transcription factor-4, exons 6-11
8358	18235	28483	8.47		AJ243213.1	TN	Homo sapiens partial 5-HT4 receptor gene, exons 2 to 5
8396	18272		3.23		BF700092.1	EST_HUMAN	602127577F1 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:4284403 5'
8482	18355		1.78		BF700092.1	EST_HUMAN	602127577F1 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:4284403 5'
3186	13111	22915	1.48	6.7E-01	Q9WTJ2	SWISSPROT	PUTATIVE TRANSCRIPTION FACTOR OVO-LIKE 1 (MOVO1) (MOVO1A)
3460	13376		2.43	5.7E-01	AB033503.1	TN	Populus euramericana peacs-2 mRNA for 1-aminocyclopropane-1-carboxylate synthase, complete cds
3829	13741	23533	1.65	6.7E-01	AF011581.1	LN	Homo sapiens T cell receptor beta chain (BV6S772-BJ1S1) mRNA, partial cds
5201	15064	24827	2.24	5.7E-01	U78517.1	IN	Rattus norvegicus cAMP-regulated guanine nucieotide exchange factor II (cAMP-GEFII) mRNA, partial cds
5836	15742	25854	3.72	5.7E-01	BF035413.1	EST_HUMAN	601454962F1 NIH_MGC_66 Homo sapiens cDNA clone IMAGE:3858590 5'
6100	15110		1.41	5.7E-01	AL111440.1	N _T	Botrytis cinerea strain T4 cDNA library under conditions of nitrogen deprivation
6564	1	26603	2.12	5.7E-01		SWISSPROT	PYRROLINE-5-CARBOXYLATE REDUCTASE (P5CR) (P5C REDUCTASE)
9124			1.68			EST_HUMAN	MR3-HT0736-180700-003-a02 HT0736 Homo sapiens cDNA
3318			1.21			NT	Homo sepiens mRNA for KIAA0740 protein, partial cds
3318	13239	23044	1.21	5.6E-01	AB018283.2	NT	Homo sapiens mRNA for KIAA0740 protein, partial cds
3808	13720		0.83		.2	NT	Arabidopsis thaliana DNA chromosome 4, contigifragment No. 13
4146	14046	23818	0.83	5.6E-01	D83135.1	NT	Chicken TBP gene, exon8, complete ods
7111	16988	27180	4.18	5.6E-01	AV684703.1	EST_HUMAN	AV684703 GKC Homo sapiens cDNA clone GKCFSF05 5'
7111	16988	27181	4.18		AV684703.1	EST_HUMAN	AV684703 GKC Homo saplens cDNA clone GKCFSF05 5'
9023	18817		2.46	∵ 5.6E-01	BE888280.1	EST HUMAN	601514007F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3915457 5'
							ng75g10.s1 NCI_CGAP_Pr6 Homo saplens cDNA clone IMAGE:940674 similar to contains element PTR7
9137	ı	28794	1.73	5.65-01	AA493535.1	EST HUMAN	repairve element;
470A	ļ		1.32			DATE INC.	THE TAIL IN THE TOTAL OF THE TAIL OF THE T
9919	19392		3.09	5.6E-01	BF573829.1	EST_HUMAN	60Z13Z0Z8F1 NIH_MGC_81 Homo sapiens cDNA clone IMAGE:4Z71334 5

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
1183	11103	20949	1.09	5.5E-01	8393912 NT	NT	Rattus norvegicus Propionyl Coenzyme A carboxylase, beta polypeptide (Pccb), mRNA
2668	12533	22423	2.88	5.5E-01	P03341	SWISSPROT	GAG POLYPROTEIN (CONTAINS: INNER COAT PROTEIN P12; CORE PROTEIN P15; CORE SHELL PROTEIN P30; NUCLEOPROTEIN P10]
							GAG POLYPROTEIN (CONTAINS: INNER COAT PROTEIN P12; CORE PROTEIN P15; CORE SHELL
8907	72033	١	j	5.55-01	705541	SWISSPROI	TAULEIN PSO, NOCKEOPRO LEIN P10
8887 78888		22608		5.5€-01		LN	Homo septens superfuller viralicidic activity 2 (5) cerevisiae homolog)-like (SKIV2L), mRNA
3027			1.39	5.5E-01	H46219.1	EST_HUMAN	yo18a10.s1 Soares adult brain N2b5HB55Y Homo sapiens cDNA clone IMAGE:178288 3'
3196	13121	97872	3.1	6.6E-01	AF227240.1	NT	Rabbit oral papillomavirus, complete genome
3634	13548	23335	1.29	5.5E-01	P48755	SWISSPROT	FOS-RELATED ANTIGEN-1
139	10113	19933	3.74	5.4E-01	7857266 NT	NT	Homo sapiens KIAA0929 protein Msx2 Interacting nuclear target (MINT) homolog (KIAA0929), mRNA
139	10113	19934	3.74	5.4E-01	7657266 NT	LN	Homo sapiens KIAA0929 protein Msx2 Interacting nuclear target (MINT) homolog (KIAA0929), mRNA
					_		Pseudomonas syringae pv. tomato strain DC3000 AvrE (avrE), HrpW (hrpW), and GstA (gstA) genes,
570	10509	20316	261	5.4E-01	AF232006.1	Ę	complete cds; and unknown genes
670	10500			5.4E.04	AF232008 4	LN.	Pseudomonas syringae pv. tomato strain DC3000 AvrE (avrE), HrpW (hrpW), and GstA (gstA) genes, complete cds: and unknown genes
1250	11157	21006	224	8.4E-01	AW896087 1	EST HUMAN	QV4-NN0040-070400-160-c04 NN0040 Homo sapiens cDNA
2059	11949			5.4E-01 A	AE002247.2	N P	Chlamydophila pneumoniae AR39, section 74 of 94 of the complete genome
2208	12095	21998		5.4E-01 A	AJ276682.1	Ę	Drosophila melanogaster mRNA for 15,15' beta carotene dioxygenase (beta-diox gene)
7738	17588			5.4E-01	5.4E-01 BF572536.1	EST HUMAN	602076545F1 NIH_MGC_62 Homo sapiens cDNA clone IMAGE:4243690 5'
8427	18301	28557		5.4E-01	P36858	SWISSPROT	NITRATE REDUCTASE (NADPH) (NR)
8892	18702		4.51	5.4E-01	Q60675	SWISSPROT	LAMININ ALPHA-2 CHAIN PRECURSOR (LAMININ M CHAIN) (MEROSIN HEAVY CHAIN)
8892	18702		4.51	6.4E-01	Q60676	SWISSPROT	LAMININ ALPHA-2 CHAIN PRECURSOR (LAMININ M CHAIN) (MEROSIN HEAVY CHAIN)
2000	4882		446	10 UV	E 4E 04 A 1960209 4	NOW IN LOS	wi37g04.x1 NCL_CGAP_Utf Homo sepiens cDNA clone IMAGE:2427126 3' similar to gb:M13452 LAMIN A
	7		2		Jinggago.	NUMBER OF THE PARTY OF THE PART	Viscon and Viscon III and an anadole and and the food of the control of the contr
							Fromo septens FiLA class III region containing tenescen A (tenescen-A) gene, peruei ods, oyocchrome F460 Z1- hydroxylase (CYP21B), complement component ©4 (C4B) G11, helicase (SKI2W), RD, complement factor B
505	10447	20260	2.02	5.3E-01	AF019413.1	Z	(Bf), and complement component C2 (C2) genes,>
2083	11982	71877	16.0	5.3E-01	AF113919.1	F	Brassica cleracea var. capitata phospholipase D2((PLD2) gene, complete ods
2093	11982	87812	0.91	5.3E-01	AF113919.1	NT	Brassica oleracea var. capitata phospholipase D2 (PLD2) gene, complete ods
2755	12617	22508	8.24	5.3E-01		NT	Homo sapiens protein tyrosine phosphatase, receptor-type, zeta polypeptide 1 (PTPRZ1) mRNA
2765	12817	22509	8.24	5.3E-01		된	Homo sepiens protein tyrosine phosphatase, receptor-type, zeta polypeptide 1 (PTPRZ1) mRNA
3205	1			5.3E-01	AF087658.1	<u>F</u>	Homo sepiens secreted C-type lectin precursor (LSLCL) gene, complete cds

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Table 4
Single Exon Probes Expressed in Heart

Top Hit Descriptor	Mycoplasma genitalium section 9 of 51 of the complete genome	zu42h12.y5 Soares overy tumor NbHOT Homo saplens cDNA clone IMAGE:740711 5'	zu42h12.y5 Soares ovary tumor NbHOT Homo sapiens cDNA clone IMAGE:740711 5'	7e73c12.x1 NCI_CGAP_Pr28 Homo sepiens cIDNA clone IMAGE:3288118 3' similar to gb:J02783 PROTEIN DISULFIDE ISOMERASE PRECURSOR (HUMAN);	7e73c12.x1 NCI_CGAP_Pr28 Homo sapiens cIDNA clone IMAGE:3288118 3' similar to gb:J02783	PROTEIN DISCLETURE SOMETHER PRECORSOR (HOMAN);	Roridula gorgonias ribulose 1,5-bisphosphate carboxylase (rbcL) gene, partial cds; chloroplast gene for chloroplast product	601339867F1 NIH_MGC_53 Homo sapiens cDNA clone IMAGE:3682168 6'	og30e05.s1 NCI_CGAP_Br7 Homo sapiens cDNA clone IMAGE:1441376 3' similar to gb:J02611 APOLIPOPROTEIN D PREGURSOR (HUMAN);	Drosophila melanogaster helix-loop-helix mRNA, complete cds	NUCLEAR FACTOR OF ACTIVATED T CELLS 5 (T CELL TRANSCRIPTION FACTOR NFAT5) (NF-AT5) (REL DOMAIN-CONTAINING TRANSCRIPTION FACTOR NFAT5)	Homo sapiens phospholipid scramblase 1 gene, complete cds	Homo sapiens chromosome 21 segment HS21C085	Homo sapiens mRNA for KIAA0740 protein, partial cds	Chlamydophila abortus strain S26/3 POMP91A and POMP90A precursor, genes, complete cds	Azotobacter vinelandii icd gene for isocitrate dehydrogenase, complete cds	Botrytis cinerea strain T4 cDNA library under conditions of nitrogen deprivation	am77g05.s1 Stratagene schizo brain S11 Homo sapiens cDNA clone IMAGE:1616504 3'	Medicego sativa chloroplast malate dehydrogenase precursor (p1mdh) mRNA, nuclear gene encoding chloroplast protein, complete cds	Homo sapiens chromosome Xq28 melanoma antigen family A2a (MAGEA2A), melanoma antigen family A12	(MAGEA12), melanoma entigen family A2b (MAGEA2B), melanoma antigen family A3 (MAGEA3), celtractin {CALT}. NAD(P)H dehydrogenase-like protein (NSDHL). and LI>	Mus musculus vanilloid receptor-like protein 1 (Vri1), mRNA	Homo sapiens PELOTA (PELOTA) gane, complete cds	UI-H-BI1-acp-a-08-0-UI.s1 NCI_CGAP_Sub3 Homo saplens cDNA clone IMAGE:2714799 3'	RETINOIC ACID RECEPTOR GAMMA (RAR-GAMMA) (RETINOIC ACID RECEPTOR DELTA) (RAR-	Human adrenodoxin reductase gene, exons 3 to 12	Polyangium vitellinum (strain Pl vt1) 165 rRNA gene	Condition of the control of the cont
Top Hit Database Source	NT	EST_HUMAN	EST_HUMAN	EST HUMAN	1444	ES HOMAN	5	EST_HUMAN	EST_HUMAN	뒫	SWISSPROT	Ę	Z	ᅜ	NT	뒫	Ę	EST_HUMAN	F		Ę	뒫	뒫	EST_HUMAN	TODOOLAGO	SWISSING!	5	-
Top Hit Acession No.	J39687.1	5.3E-01 AI820921.1	41820921.1	, 3E645620.1		3E645620.1	.01950.2	3E566291.1	5.3E-01 AA916053.1	20770.1	Q9WV30	AF224492.1	4L163285.2	5.2E-01 AB018283.2	J65942.1	D73443.1	5.2E-01 AL116780.1	4A984165.1	AF020289.1		J82671.2	7106444 NT	AF143952.2	4W137068.1		M58509 1	,	47500044.1
Most Similar (Top) Hit BLAST E Value	5.3E-01	5.3E-01	6.3E-01	5.3E-01	100	5.3E-01 B	5.3E-01	5.3E-01	5.3E-01	5.2E-01	5.2E-01	6.2E-01	5.2E-01	5.2E-01	6.2E-01	5.2E-01	5.2E-01	5.2E-01	5.2E-01		5.2F-01	5.25-01	6.2E-01	5.2E-01	100	5.4F.01	5.4E-04	0.1771
Expression Signal	1.29	1.75	1.76	1.91		1.91	2.15	5.62	246	11.31	7.69	2.83	5.04	2.75	1.9	1.22	1.76	1.9	1.13		1.08	1.29	1.28	2.46	9	2,43	3.28	3.441
ORF SEQ ID NO:			26089	25447		25448		28946		20568	20904	20830		21886				23114			23287		27837	25221			20384	
Exon SEQ ID NO:	14017	15270	15270	15387	L	12287	17038		19551	10728	11061		l_	11988	13008	13124	ļ	13314	13494		13496	1_	17611	19260		10537	L	╛
Probe SEQ ID NO:	4117	5350	5350	5467		5467	7161	8846	9015	799	1148	1174	1843	2089	3081	3199	3359	3397	3580		3582	4949	7761	9719		9887	3	733

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Single Exon Probes Expressed in Heart

						,	
Probe SEQ ID	Exon SEQ ID	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E	Top Hit Acession No.	Top Hit Database Sourse	Top Hit Descriptor
į	<u> </u>			Vatue			
632	10569	20382	3.28	5.1E-01	AJ233944.1	TN	Polyangium vitellinum (strain PI vt1) 16S rRNA gene
1632	11536		1.06	5.1E-01	X87885.1	IN.	R.norvegicus mRNA for mammalian fusca protein
1977	11870		1.44	5.1E-01	BF683095.1	EST_HUMAN	602139319F1 NIH_MGC_46 Homo sapiens cDNA clone IMAGE:4298117 5
3984	13891	23667	3.84	5.1E-01	AI858495.1	EST_HUMAN	w89b/12.x1 NCI_CGAP_Ut1 Homo sapiens cDNA clone IMAGE:2427263 3'
4094		23771	2.86	5.1E-01	P96380	SWISSPROT	TRANSCRIPTION-REPAIR COUPLING FACTIOR (TRCF)
6133	15980	26116	1.57	5.1E-01	R80873.1	EST_HUMAN	y94a09.s1 Soares placenta Nb2HP Homo sapiens cDNA clone IMAGE:1468723'
7584	17435	27650	4.54	5.1E-01	J05412.1	NT	Human regenerating protein (reg) gene, complete cds
7586	17437	27653	3.57	5.1E-01	W22302.1	EST_HUMAN	65B1 Human retina cDNA Tsp509I-cleaved sublibrary Homo sapiens cDNA not directional
9230	19478		3.62	5.1E-01	BF030207.1	EST_HUMAN	601556863F1 NIH_MGC_58 Homo sapiens cDNA clone IMAGE:3826767 5'
0473	19100		203	8 15.01	PF430082 1	NAMILH TRE	nac51f10.x1 NCI_CGAP_Brn23 Home sapiens cDNA clone IMAGE:3406218 3' similar to contains element TAR1 repetitive element
7806		24874	1 97	70 II O	ш_	LN	Homo sanjans metmalolic segregation Increased 2.ilke 9 (PMS2) 9) mRNA
2007		24872	137	2.0F.04		5 5	Homo sapians postmeiotic segregation increased 2-like 9 (PMS21.9) mRNA
7007		71017	5:	2.00		INI	
							Buchnera aphidicola genomic fragment contatring (chaperone Hsp60) groEL, DNA biosynthesis initiating protein (dnaA), ATP operon (atpCDGAHFEB), and putative chromosome replication protein (gldA) genes,
2097	11988	21882	1.32	5.0E-01	AF008210.1	님	complete cds; and termination factor Rho (rho) gene>
							Buchnera aphidicola genomic fragment containing (chaperone Hsp60) groEL, DNA biosynthesis initiating
							protein (dnaA), ATP operon (atpCDGAHFEB), and putative chromosome replication protein (gldA) genes,
2097			1.32	6.0E-01	AF008210.1	Ę	complete cds; and termination factor Rho (rho) gene>
3688	13601	23388	0.84	5.0E-01	U55574.1	TN.	Mus musculus anti-DNA immunoglobulin light chain IgM mRNA, antibody 363p.138, partial cds
3797	13709		3.44	5.0E-01	AB033010.1	ŊŢ	Homo sapiens mRNA for KIAA1184 protein, partial cds
6961	16839		5.62	5.0E-01	M92304.1	NT	Xenopus laevis smooth muscle beta-tropomyosin mRNA, complete cds
7442	16455	26644	3.61	5.0E-01	BF317212.1	EST_HUMAN	601803871F1 NIH_MGC_19 Homo saplens cDNA clone IMAGE:4136332 5
							GLYCOGEN DEBRANCHING ENZYME (GLYCOGEN DEBRANCHER) (INCLUDES: 4-ALPHA-
7550	17401	27814	2.26	5.0E-01	P35573	SWISSPROT	GEUGANO (RANSTERASE (CEGO-1,1-1,1-16-10-14) (DEXTRING BENGE), AMITEO-1,9-16-10-003-10-3-16-10-10-14-15-10-10-10-10-10-10-10-10-10-10-10-10-10-
	1				1		GLYCOGEN DEBRANCHING ENZYME (GLYCOGEN DEBRANCHER) [INCLUDES: 4-ALPHA-
							GLUCANOTRANSFERASE (OLIGO-1,4-1,4-GLUCANTRANSFERASE); AMYLO-1,8-GLUCOSIDASE
7550	17401	27615		5.0E-01	P35573	SWISSPROT	(DEXTRIN 6-ALPHA-D-GLUCOSIDASE)]
7941			1.27	5.0E-01	BE869218.1	EST_HUMAN	601445024F1 NIH_MGC_65 Homo sapiens cDNA clone IMAGE:3849438 5'
8003	18806	28099	. 9.32	5.0E-01	1	NT	Oryzlas latipes gene for membrane guanyly cyclase OIGC1, complete cds
9170	18913		2.26	5.0E-01	AF029215.1	ᅜ	Mus musculus MRC OX-2 antigen homolog gene, exons 2-5, and complete cds
9828			2.02	5.0E-01	AL163302.2	F	Homo sapiens chromosome 21 segment HS21C102
6986	19358		2.94	5.0E-01	013961	SWISSPROT	NUCLEAR ENVELOPE PROTEIN CUT11

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Single Exon Probes Expressed in Heart

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Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Vätue	Top Hit Acesslan No.	Top Hit Database Source	Top Hit Descriptor
5414	15334	25384	3.29	4.6E-01	AI247679.1	EST_HUMAN	qh59h02x1 Soares_fetal_iiver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:1849011 3' similar to TR:016338 016338 BUTYROPHILIN.;
5414	16334		3.29	4.6E-01	AI247679.1	EST_HUMAN	qh59h02x1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:1849011 3' similar to TR:O15338 O15338 BUTYROPHILIN.;
5418		25393		4.6E-01	P20050	SWISSPROT	MEIOSIS SPECIFIC PROTEIN HOP1
6057	16040		1.47	4.6E-01	U62332.1	IN	Emericella nidulans NEMPA (nempA) gene, mitochondrial gene encoding putative mitochondrial protein, complete cds
5908	76040			185 04	1 180000 4	FIN	Emericella nidulans NEMPA (nempA) gene, mit/chondrial gene encoding putative mitochondrial protein,
6843	┸	26916		4.6E-01	<u> </u>	EST HUMAN	602130953F1 NIH _MGC _56 Homo saplens cDNA clone IMAGE:4287828 5'
7363	17230	27430	26.48	4.6E-01	P55202	SWISSPROT	ATRIAL NATRIURETIC PEPTIDE RECEPTOR. B PRECURSOR (ANP-B) (ANPRB) (GC-B) (GUANYLATE CYCLASE)
7363	1			4 65-01		SWISSPROT	ATRIAL NATRIURETIC PEPTIDE RECEPTOR B PRECURSOR (ANP-B) (ANPRB) (GC-B) (GUANYLATE CYCLASE)
7730	1			4.6E-01	. ~	EST HUMAN	wg73e12.x1 Soares_NSF_F8_9W_OT_PA_P_\$1 Homo sapiens cDNA clone IMAGE:2370766 3'
7730					AI915634.1	EST_HUMAN	wg73e12.x1 Soares_NSF_F6_9W_OT_PA_P_S1 Homo sapiens cDNA clone IMAGE:2370766 3'
8328	18236		2.86		ш	SWISSPROT	PUTATIVE VITELLOGENIN RECEPTOR PRECURSOR (YL)
8368				4.6E-01	w	EST_HUMAN	IL6-HT0730-100500-075-g05 HT0730 Homo sapiens cDNA
8368				4.6E-01	ш	EST_HUMAN	L5-HT0730-100500-075-g05 HT0730 Homo sapiens cDNA
8766	17915	28160	5,45	4.6E-01	AF019369.1	NT	Human thiopurine methyltransferase (TPMT) gene, exon 10 and complete cds
8766	17915	28161	5.45	4.6E-01	AF019369.1	NT	Human thiopurine methyltransferase (TPMT) gene, exon 10 and complete cds
9313	19005		1.43	4.6E-01	D53316.1	EST_HUMAN	HUM105F03B Clontech human fetal brain polyA÷ mRNA (#6535) Homo sapiens cDNA clone GEN-105F03 5'
9978	19491		2.21	4.6E-01	AF120134.1	N	Linanthus jamauensis maturase (matK) gene, chioropiast gene encoding chloropiast protein, partial cds
1869	11765	21639	1.73	4.5E-01	AE001931.1	TN	Delnococcus radiodurans R1 section 68 of 229 of the complete chromosome 1
1869	11765			4.6E-01		NT	Deinococcus radiodurans R1 section 68 of 229 of the complete chromosome 1
2841	12769	22558	4.87	4.5E-01	AA677086.1	EST_HUMAN	455402.s1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens oDNA clone IMAGE:4541793'
3276	13196	22996	3.97	4.5E-01	Q05783	SWISSPROT	BASEMENT MEMBRANE-SPECIFIC HEPARAN SULFATE PROTEOGLYCAN CORE PROTEIN PRECURSOR (HSPG) (PERLECAN) (PLC)
3331	L	L	1.05	4.5E-01	AF126378.1	TN	Mus musculus DNA polymerase epsilon catalytic subunit (Pole) gene, exons 2 through 12
3942			1.41	4.5E-01		SWISSPROT	COLLAGEN ALPHA 5(IV) CHAIN
3982		23665		4.5E-01		EST_HUMAN	as96e09.x1 Barstead aorta HPLRB6 Homo sapiens cDNA clone IMAGE:2353480 3'
4085	15079		4.02		AW873495.1	EST_HUMAN	hod0g02.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:3041810 3

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Single Exon Probes Expressed in Heart

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	Top Hit Descriptor	601657225R1 NIH_MGC_67 Homo sepiens cDNA clone IMAGE:3868023 3'	QV2-PT0012-140100-031-c09 PT0012 Homo sepiens cDNA	COAT PROTEIN	W32e02x1 NCI_CGAP_Ut1 Homo sapiens cDNA clone IMAGE:2426618 3' similar to TR:Q92923 Q92923 SWIISNF COMPLEX 170 KDA SUBUNIT.	tz56g11.x1 NCI_CGAP_Ov35 Homo saplens cDNA clone IMAGE:2292644 3'	Homo sapiens hypothetical protein DKFZp547G183 (DKFZp547G183), mRNA	EST02531 Fetal brain, Stratagene (cat#936206) Homo sapiens cDNA clone HFBCY17	EST02531 Fetal brain, Stratagene (cat#936206) Homo sapiens cDNA clone HFBCY17	xo14h01.x1 NCI_CGAP_Uß Homo sepiens cDNA clone IMAGE:2703985 3' sImiter to SW:INT6_MOUSE Q64252 VIRAL INTEGRATION SITE PROTEIN INT-6. [1];	601449201F1 NIH_MGC_65 Homo seplens cDNA clone IMAGE:3852961 5'	602035275F1 NCI_CGAP_Brn64 Homo sapiens cDNA clone IMAGE:4183290 5'	Homo saplens testis-specific kinase 2 (TESK2), mRNA	Entamoeba histolytica diaphanous protein (dia) gene, partial cds	Mus musculus integral membrane-associated protein 1 (Itmap1), mRNA	VASCULAR ENDOTHELIAL GROWTH FACTOR B PRECURSOR (VEGF-B) (VEGF RELATED FACTOR)	Rattus norvedicus SvnGAP-b mRNA, complete cds	Rattus norvegicus SynGAP-b mRNA, complete cits	7j91d02.y1 NCI_CGAP_Br16 Homo sapiens cDNA clone IMAGE:3393795 5'	801237139F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3609393 5'	MR0-HT0078-131299-007-g05 HT0078 Homo sapiens cDNA	HISTIDINE-RICH GLYCOPROTEIN PRECURSOR	HISTIDINE-RICH GLYCOPROTEIN PRECURSOR	mucin (rats, Spregue-Dawley, sulfur-dioxide-treated tracheal epithelium, mRNA Partial, 390 nt)	AV720408 GLC Homo sepiens cDNA clone GLCCSC12 5'	qi62h11.x1 NCI_CGAP_Brn25 Homo sapiens cDNA clone IMAGE:1861125 3' similar to TR:Q29168 Q29168 UNKNOWN PROTEIN;	qi82h11.x1 NCI_CGAP_Brn25 Homo sapiens cDNA clone IMAGE:1861125 3' sImilar to TR:Q29168 Q29168 UNKNOWN PROTEIN	xx27e08.x1 NCI_CGAP_Co18 Homo sapiens cDNA clone IMAGE:2585510 3' similar to TR:095154 095154 AFLATOXIN B1-ALDEHYDE REDUCTASE. ;	S.tuberosum mRNA for Induced stolon tip protein (partial)
	Top Hit Database Source	EST_HUMAN	EST_HUMAN	SWISSPROT	EST HUMAN	EST HUMAN	N	EST_HUMAN	EST_HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	TN	TN	NT	SWISSPROT	ŀ	NT.	EST_HUMAN	EST_HUMAN	EST_HUMAN	SWISSPROT	SWISSPROT	NT	EST_HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	ŢN
	Top Hit Acession No.	BE983445.2	AW608814.1		A1858849.1	AI648596.1	11444786 NT	M86006.1	M86006.1	AW591271.1		BF337531.1	11422099 NT	AF238234.2	6680503 NT	P49765	AF058790 1	AF058790.1	4.4E-01 BF056726.1	BE378707.1	BE141396.1	P04929	P04929	S65019.1	AV720408.1	A[198413.1	AI198413.1	AW080795.1	Z11679.1
	Most Similar (Top) Hit BLAST E Value	4.5E-01	4.5€-01	4.SE-01	4.5E.01	4.50-04	4.5E-01	4.5E-01	4.5€-01	4.5E-01	4.6E-01	4.5E-01	4.5E-01	4.5E-01	4.4E-01	4 4F-01	4 4F-03	4.4E-01	4.4E-01	4.4E-01	4.4E-01	4.4E-01	4.4E-01	4.4E-01	4.4E-01	4.4E-01	4.4E-01	4.4E-01	4.4E-01
	Expression Signal	0.94	1.45	1.47	2.49	3.14	1.54	25.09	25.09	2.42	2.26	1.44	2.15	1.83	2.47	9,50	80 0	0.99	1.87	1.33	0.93	1.99	1.99	1.3	1.98	1.6	91	1.79	10.84
	ORF SEQ ID NO:		25375	L		26970		28082	28083					24987		22118			22398			25050	25051						
	Exan SEQ ID NO:	14742	ı	1	ļ	Ι_	L	17841	17841	18117	1		19283	١.	<u> </u>	L	1.			_	L	15246	15246	15409	_			1	ı
	Probe SEQ ID NO:	4862	5408	5983	6488	6897	7100	7991	7991	8237	9036	9694	9763	9974	1991	2240	3274	3274	3277	4141	4924	5326	5326	5490	5499	8624	FR24	5782	6582

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Single Exoll Flobes Explessed III near	Top Hit Descriptor	ZINC FINGER X-CHROMOSOMAL PROTEIN	qo39f09.x1 NCI_CGAP_Lu5 Homo sapiens cDNA clone IMAGE:1910921 3'	GLYCOPROTEIN B PRECURSOR (GLYCOPROTEIN 14)	TYROSINE-PROTEIN KINASE RECEPTOR TIE-1 PRECURSOR	beta -HKA=H,K-ATPase beta-subunit [rats, Genomic, 8983 nt, segment 2 of 2]	beta -HKA=H,K-ATPase beta-subunit [rats, Genomic, 8983 nt, segment 2 of 2]	Mus musculus sodium channel, type X, alpha polypeptide (Scn10a), mRNA	Homo sapiens chromosome 21 segment HS210082	Autographa californica nucleopolyhedrovirus, complete genome	UV EXCISION REPAIR PROTEIN PROTEIN RAD23 HOMOLOG A (HHR23A)	Callithrix Jacchus MW/LW opsin gene, upstream flanking region	Callithrix jacchus MW/LW opsin gene, upstream flanking region	QV4-SN0024-200400-183-b01 SN0024 Homo sepiens cDNA	CM2-DT0003-010200-077-c01 DT0003 Homo sepiens cDNA	MRo-BN0070-270300-008-g04 BN0070 Homo septens cDNA	Aquifex aeolicus section 30 of 109 of the complete genome	Human somatostatin I gene and flanks	Callithrix Jacchus MW/LW opsin gene, upstream flanking region	Callithrix Jacchus MW/LW opsin gene, upstream flanking region	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 14	Xestia c-nigrum granulovirus, complete genome	Salmiri sclureus olfactory receptor (SSC186) gene, partial cds	Coturnix coturnix japonica ifnG gene	DNA GYRASE SUBUNIT B	602023134F1 NCI_CGAP_Bm67 Homo sapiens cDNA clone IMAGE:4158298 5'	Methanococcus voltae flagella-related protein C-I (flaC-flal) genes, complete cds	hh74e10.y1 NCI_CGAP_GU1 Homo sapiens cDNA clone IMAGE:2968554 5'	hh74e10.y1 NCI_CGAP_GU1 Homo septens cDNA clone IMAGE:2988554 5'	xn63e05.x1 Soares_NHCeC_cervical_tumor Homb sepiens cDNA clone IMAGE:2698400 3' similar to	TR:000189 000189 MU-ADAPTIN-RELATED PROTEIN 2.;	Equus caballus microsatellite LEX027	RC3-BN0034-290200-013-c12 BN0034 Homo sapiens cDNA	RC3-BN0034-290200-013-o12 BN0034 Homo sapiens oDNA	Streptomyces coelicolor whilh gene
שום באחוו היום	Top Hit Database Source	SWISSPROT	EST_HUMAN	SWISSPROT	SWISSPROT	TN	TN	NT	NT	TN	SWISSPROT	NT	IN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	IN	된	IN	NT	IN	NT	TN	SWISSPROT	EST HUMAN	IN	EST_HUMAN	EST_HUMAN		EST HUMAN	Ā	EST_HUMAN	EST_HUMAN	된
iio	Top Hit Acession No.	062836	Al268650.1	P28922	P35590	S76404.1	S76404.1	6677874 NT	AL183282.2	9627742 NT	P54725	AF155218.1	AF155218.1	AW866550.1		AW889477.1	AE000698.1	J00306.1	AF155218:1	AF156218.1	AL161502.2	9635250 NT	AF179825.1	AJ001678.1		.1	U97040.1		AW630048.1		AW170559.1	AF075629.1	AW993658.1	AW993658.1	AJ003022.1
	Most Similar (Top) Hit BLAST E Vatue		4.4E-01	4.4E-01	4.4E-01	4.4E-01	4.4E-01	4.4E-01	4.4E-01	4.4E-01		4.3E-01	4.3E-01	4.3E-01	4.3E-01	4.3E-01	4.3E-01	4.3E-01	4.3E-01	4.3E-01	4.3E-01	4.3E-01	4.3E-01	4.3E-01		4.3E-01	4.3E-01	4.3E-01	4.3E-01		4.3E-01				4.3E-01
	Expression Signal	1.29	1.89	2.45	4.67	1.33	1.33	2.29	3.29	5.34	1.75	1.98	1.98	1.1	0.98	0.93	1.53	1.15	1.02	1.02	1.11	1.06	3.04	3.86	4.01	2.54	2.83	1.65	1.65		1.27	2.64	1.98	1.98	2.02
	ORF SEQ ID NO:		27798		27903			25329		25224		20178	20179	. 21350		22742		23737	20178	20179				26064			-	27874	27875					28674	
	Exon SEQ ID NO:	17222	17573	17574	17663	17785	17785	18996	19863	19267	19326	10351	10351	11489	12771	12950	13145	13981	10351	10351	14755	14929	15514	15933	15996	16255	16787	17459	17459		17722	16049	18409	1	19390
	Probe SEQ ID NO:	7354	7723	7724	7813	7935	7935	9538	9308	9731	9824	405	405	1585	2843	3022	3221	4059	4308	4308	4876	5059	2600	6028	6102	සමය	6069	7608	7608		7872	8302	8537	8537	9916

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	. Top Hit Descriptor
1335	12691	21089	1.04	4.2E-01	Q39102	SWISSPROT	CELL DIVISION PROTEIN FTSH HOMOLOG PRECURSOR
1904	11800		0.89	4.2E-01	4.2E-01 AA761653.1	EST_HUMAN	nz24a09.s1 NCI_CGAP_GCB1 Homo sapiens ci0NA clone IMAGE:1288696 3'
3558	13472	23263	4.78	4.2E-01	E003947.1	IN	Xylella fastidiosa, section 93 of 229 of the complete genome
3585			1.09	4.2E-01	AI280338.1	EST_HUMAN	ql94b01.x1 Soares_NhHMPu_S1 Homo saplens cDNA clone IMAGE:187994531
3803	13715		0.96	4.2E-01	AW835527.1	EST_HUMAN	QV0-LT0015-180200-127-h01 LT0015 Hamo sapiens cDNA
3900	13810		1	4.2E-01	4.2E-01 Q04886	SWISSPROT	SOX-8 PROTEIN
4545	14438		3.63	4.2年-01	BE073574.1	EST HUMAN	RC5-BT0559-020300-013-E06 BT0559 Homo saplens cDNA
4599	14487	24273	4.96		AA534093.1	EST_HUMAN	nj89h01.s1 NCJ_CGAP_Pr10 Homo saplens cDNA clone IMAGE:997777 similær to gb:M33600 HLA CLASS II HISTOCOMPATIBILITY ANTIGEN, DR-1 BETA CHAIN (HUMAN);
4671	14557	24350		4.2E-01 F	R13467.1	EST_HUMAN	y777e01.r1 Soares infant brain 1NIB Homo sapiens cDNA clone IMAGE:28278 5'
5503	15422	26484	1.5	4.2E-01	BF242055.1	EST_HUMAN	601879721F1 NIH_MGC_55 Homo saplens cDNA clone IMAGE:4108493 5'
5533	15450	25518	1.58	4.2至-01	AW854162.1	EST_HUMAN	RC3-CT0254-060400-029-g04 CT0254 Homo sapiens cDNA
6151	16024		9.1	4.2E-01	AU158472.1	EST_HUMAN	AU158472 PLACE2 Homo saplens cDNA clone PLACE2000470 3'
6151	16024			4.2E-01	3U158472.1	EST_HUMAN	AU158472 PLACE2 Homo sapiens cDNA done PLACE2000470 3
6184	19461			4.2E-01		NT	Brca1=breast cancer gene [rats, WF, spleen, Genomic, 419 nt, segment 2 of 2]
6217	16083	26233	5.91	4.2E-01	2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 47
6663	16543	26739	2.1	4.2是-01	4.2E-01 AW957448.1	EST_HUMAN	EST369413 MAGE resequences, MAGE Homo sapiens cDNA
6663	16543	26740	2.1	4.2E-01	4.2E-01 AW957448.1	EST_HUMAN	EST369413 MAGE resequences, MAGE Homo sapiens cDNA
6776	16655	26844	1.19		4758039 NT	N	Homo sapiens cytochrome c oxidase subunit VIc (COXBC), nuclear gene encoding mitochondrial protein, mRNA
7992			1.48	4.2E-01	\W863666.1	EST_HUMAN	MR3-SN0010-280300-103-h07 SN0010 Homo sapiens cDNA
8400	18276		2.2		\B023489.1	NT	Oryzias latipes OIGC7 mRNA for membrane guanyly cyclase, complete cds
6698	18517	28789	2.04	4.2E-01	BE966485.2	EST_HUMAN	601660352R1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3906085 3'
9863	19353				4.2E-01 AV731815.1	EST HUMAN	AV731815 HTF Homo sapiens cDNA clone HTFBHH05 5'
1078	10994		1.88	4.1E-01	ч.	EST_HUMAN	RC-BT091-210199-142 BT091 Homo sapiens cDNA
1087					I Q I	EST_HUMAN	AV705243 ADB Homo sapiens cDNA clone ADBAHF08 5'
1087	11003	20845	24	4.15-01	AV705243.1	EST_HUMAN	AV705243 ADB Homo saplens cDNA clone ADBAHF08 5'
1592				4.1E-01	AI905949.1	EST_HUMAN	PM-BT103-270499-684 BT103 Homo sapiens cDNA
2678					7705283	NT	Homo sapiens anaphase-promoting complex subunit 7 (APC7), mRNA
2910		22633			AL161536.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 36
2910					AL161536.2	L/	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 36
3263						EST_HUMAN	oj94b08.s1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:1505943 3'
4177	i	23852			AJ249207.1	Į.	Rhodococcus sp. AD45 IsoG, IsoH, IsoI, IsoA, IsoA, IsoB, IsoC, IsoD, IsoE and IsoF genes
4208	14107		1.07	4.1E-01	AA909257.1	EST_HUMAN	am33d02.s1 Scares_NFL_T_GBC_S1 Hama saplens cDNA clone IMAGE:1542819 3

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Single Exon Probes Expressed in Heart

SEQ ID SEQ ID IC NO: NO: NO: 4373 14259						
14459	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
14459		0.83	4.1E-01	R41726.1	EST_HUMAN	yg11b03.s1 Soares infant brain 1NIB Homo sapieris cDNA clone IMAGE:31814 3'
450.40	24247	1.26	4.1E-01	AV747880.1	EST_HUMAN	AV747880 NPC Homo sapiens cDNA clone NPCBDF10 5'
20.40		0.87	4.1E-01	299124.1	NT	Bacillus subtilis complete genome (section 21 of 24); from 3999281 to 4214814
15557	25650	4.13	4.1E-01	BF681393.1	EST_HUMAN	602166690F1 NIH_MGC_83 Hamo sapiens cDNA clone IMAGE:4297319 6'
16257	26418	2.91	4.1E-01	U67535.1	NT	Methanococcus jannaschii section 77 of 150 of the complete genome
16563	28757	1.3	4.1E-01	BF574604.1	EST_HUMAN	602133281F1 NIH_MGC_81 Homo sapiens cDNA clone IMAGE:4288238 5
7253 17130	27323	1.23	4.1E-01	6755521 NT	Ę	Mus musculus signaling intermediate in Toll pathway-evolutionarily conserved (Sitpec-pending), mRNA
17738		1.3	4.1E-01	AL139076.2	NT	Campylobacter jejuni NCTC11168 complete genorine; segment 3/6
8018 17868 · ·		1.58	4.1E-01		EST_HUMAN	CM2-HT0137-200999-010-e08 HT0137 Homo sapiens cDNA
8213 18097	28349	164	4.1E-01	X58700.1	NT	Zea mays ZMPMS2 gene for 19 kDa zein protein
8696 17880	28121	2.79	4.1E-01	Q09470	SWISSPROT	VOLTAGE-GATED POTASSIUM CHANNEL PRÓTEIN KV1.1 (HUKI) (HBK1)
9624 19702		1.72	4.1E-01	D87675.1	N	Homo sapiens DNA for amyloid precursor protein, complete cds
10940	20783	1.27	4.0E-01	8404656 NT	LZ.	Laqueus rubellus mitochondrion, complete genome
1316 11223	21079	0.98	4.0E-01	AF203478.1	NT	Drosophila melanogaster Dalmatian (dmt) mRNA, complete cds
1469 11374		4.06	4.0E-01	6879258 NT	NT	Mus musculus platelet derived growth factor receptor, beta polypeptide (Pdgfrb), mRNA
1960 12709	21742	1.17	4.0E-01	296933.1	NT	Ascobolus immersus masc2 gene
12709	21743	1.17	4.0E-01	Z96933.1	NT	Ascobolus immersus masc2 gene
10110	18831	1.09	4.0E-01	LN 0678490 NT	LN	Mus musculus ubiquitin-protein ligase e3 componen n-recognin (Ubr1), mRNA
12866	22665	224	4.0E-01	AL163280.2	ĽN.	Homo sapiens chromosome 21 segment HS21C080
2939 12866	22666	2.24	4.0E-01	AL163280.2	NT	Homo saplens chromosome 21 segment HS21C080
						Streptococcus pneumoniae YilC (yilC), YilD (yilD), ipenicillin-binding protein 2x (pbp2x), and undecaprenyl- phosphate-UDP-MurNAc-penlapeptide phospho-MurNAc-penlapeptide transferase (mraY) genes, complete
3637 13551	23339	1.61	4.0E-01	AF068903.1	N	spo
3755 13668	23451	3.24	4.0E-01	AJ277511.1	TN	Ovis aries partial JD2 gene for T cell receptor delta chain (TCRDJ2), exon 1
3755 13668	23452	3.24	4.0E-01		N	Ovis aries partial JD2 gene for T cell receptor delta chain (TCRDJ2), exon 1
4712 14598	-	7.34	4.0E-01		SWISSPROT	NADH-PLASTOQUINONE OXIDOREDUCTASE CHAIN 5, CHLOROPLAST
5603 15517	25596	1.31	4.0E-0:1	AW970610.1	EST_HUMAN	EST382691 MAGE resequences, MAGK Homo saplens cDNA
8959 18766		3.24	4.0E-01	L76080.1	TN.	Synechocystis sp. PCC 9413 transposase gene, complete cds
9312 19567	-	2.22	4.0E-01	AL163300.2	NT	Homo sapiens chromosome 21 segment HS21C100
9825 19327		1.33	4.0E-01	P36049	SWISSPROT	HYPOTHETICAL 49.7 KD PROTEIN IN GIN2-STE3 INTERGENIC REGION
9910 19387		1.4	4.0E-01		NT	Campylobacter jejuni NCTC11168 complete genome; segment 2/6
	20002	1.59	3.9E-01	1	EST_HUMAN	CM4-HT0136-150999-014-109 HT0136 Homo sapiens cDNA
1356 11262	21119	2.1	3.9E-01	AF206618.1	IN	Gorilla gorilla carboxyl-ester lipase (CEL) gene, complete cds

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Top Hit Descriptor	Homo sapiens mRNA for KIAA1183 protein, partial cds	H.sapiens B-myb gene	H.saplens B-myb gene	Sinorhizobium meliloti egi, syrB2, cya3 genes and orf3	7161401.x1 NCI_CGAP_Br16 Hamo sapiens cDNA clone IMAGE:3339169 3'	601563948F1 NIH_MGC_20 Hamo sapiens cDNA clane IMAGE:3833699 5'	601862362F1 NIH_MGC_53 Homo sapiens cDNA clone IMAGE:4082055 5'	xn88d04.x1 Soares_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:2701351 3' similar to TR:094821 O94821 KIAA0713 PROTEIN;	wp76a02.x1 NCI_CGAP_Bm25 Homo sapiens cDNA clone IMAGE:2467658 3' similar to SW:RFX5 HUMAN P48382 BINDING REGULATORY FACTOR.;	Human clabindin 27 gene, exons 10 and 11, and II.1 and Alu repeats	AV695974 GKC Homo sapiens cDNA clone GKCBQC11 5'	Homo sapiens proteoglycan 3 (PRG3) gene, complete cds	HOMEOBOX PROTEIN HLX1	Thermotoga maritima section 123 of 136 of the complete genome	Homo saplens protein kinase PKNbeta (pknbeta), imRNA	Xyella fastidiosa, section 16 of 229 of the complete genome	Arabidopsis thaliana putative c-myb-like transcription factor (MYB3R-3) mRNA, complete cds	Mus musculus solute carrier family 1, member 6 (SIc1a6), mRNA	Human immunodeficiency virus type 1 complete genome (isolate 98SE-MP1213)	Pleuronectes americanus aminopeptidase N (ampN) gene, partial cds		wf38b12.xl Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2357855 3'	wf38b12.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2357855 3'	Mus musculus general transcription factor II I (Gtf2i), mRNA	601074110F1 NIH_MGC_12 Homo sapiens cDNA clone IMAGE:3460154 5'	TRANSCRIPTION FACTOR SOX-10	QV3-BT0537-271299-049-e02 BT0537 Homo saplens cDNA	ta54f11.x1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA clone IMAGE:2047917 3' similar to	contains Afu repetitive element;	M.musculus gene for kallikrein-binding protein	Homo sapiens mRNA for KIAA1631 protein, partiel cds
Top Hit Database Source	IN	IN		NT	EST_HUMAN	EST_HUMAN	EST_HUMAN			Γ	EST_HUMAN /	Z	Г	N					TN	I		HUMAN	T_HUMAN			SWISSPROT	EST_HUMAN		EST_HUMAN		Ę
Top Hit Acessian No.	AB033019.1	X82032.1	X82032.1	AJ225896.1	BF592611.1	BE728667.1	BF208036.1	AW195888.1	Al937337.1	M19879.1	AV695974.1	AF304354.1		AE001811.1	7019488 NT	AE003870.1	AF214117.1	6678002 NT	AJ251057.1	AF043383.1	AL161518.2	AI807219.1	AI807219.1	6754095 NT	BE544653.1	Q04888	BE072399.1		AI374601.1	X61597.1	AB046851.1
Most Similar (Top) Hit BLAST E Value	3.9E-01	3.9E-01	3.9E-01	3.9E-01	3.9€-01	3.95-01	3.9E-01	3.9E-01		3.9E-01	3.8E-01	3.9E-01	3.9E-01	3.9E-01	3.8E-01	3.8E-01	3.8E-01	3.8E-01	3.8E-01	3.85-01	-		_	3.8E-01	3.8E-01	3.8E-01	3.8E-01	_			3.8E-01
Expression Signal	3.54	5.98	5.98	3.63	1.47	1.48	3.63	1.5	4.	2.97	222	249	238	1.26	7.44	1.22	2.44	4.41	0.92	2	77.7	67.0	16.0	0.85	0.98	1.62	4.47		3.81	4.83	3.07
ORF SEQ ID NO:	22367	22427	22428	22776	23672	24572	25611	27400	27581	27755				25328			22289	22352		22732	23160			23645	24692	25422	26023		26177		27052
Exan SEQ ID NO:	12473	12538	12538	12985	13895	14798	15528	17200	17372	17530	18080	19642	18940	18990	10128	11724	12398	12728	12899	12938	13355		13410	13752	14918	15366	15899		16036	1	16857
Probe SEQ ID NO:	2605	2673	2673	3058	3988	4919	5613	7324	7603	7680	8195	9091	8209	9288	154	1827	2524	2691	2972	3010	3438	3494	3508	3841	5048	5445	5894		6053	6438	9869

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Single Exon Probes Expressed in Heart

	00F SEQ 10 ID NO: 10 ID NO	Signa		op Hit Acessian No. No. No. 100. 100. 100. 100. 100. 100. 100. 10	Top Hit Source Source Source T HUMAN T HUMAN T HUMAN T HUMAN T HUMAN T HUMAN T HUMAN T HUMAN T HUMAN T HUMAN T HUMAN T HUMAN	Top Hit Descriptor Nex3h06.rt Sceres fetal liver spleen NNF.S Hom's septens oDNA clone IMAGE:120539 6' similar to contains Au repetitive element contains PTRS repetitive element; RCD-H10641-0408000-032-b12 H10641 Hom espiens cDNA clone IMAGE:30289 3' RCD-H10641-0408000-032-b12 H10641 H0m espiens cDNA clone IMAGE:3089 3' RCD-H10641-0408000-032-b12 H10641 H0m espiens cDNA clone IMAGE:3089 3' RCD-H10641-0408000-032-b12 H10641 H0m espiens cDNA clone IMAGE:3089 3' RCD-H10641-0408000-032-b12 H10641 H0m espiens cDNA clone 1291657 3' similar to similar to general sceptory VIRA4 (VII:44) gene, complete cds RCD-ETATE REPORT SERVIN ALEUMIN HOTERS.; RMS-DTO007-080300-1044-b20 T0007 H0m espiens cDNA clone IMAGE:3080652 5' RGS-BTA Searce Leste SMA CTE A STARM MCGe section 60 of 206 of the complete cds RGS-BTA Searce NSF F8 9W OT PA STARM CARPS A MINA Adone IMAGE:3080652 5' RGC-BTA STARM CARPS A STARM MCGE SECTION CARPS A MINA A GONE SEPTEN HIM MGC G9 Hom ospiens cDNA clone IMAGE:3080652 5' RGC-148089777 INIH MGC G9 Hom ospiens cDNA clone IMAGE:3080652 5' RGC-148089777 INIH MGC G9 Hom ospiens cDNA clone IMAGE:3080652 5' RGC-148089777 INIH MGC G9 Hom ospiens cDNA clone IMAGE:3080652 5' RGC-148089777 INIH MGC G9 Hom ospiens cDNA clone IMAGE:3080652 5' RGD-148089777 INIH MGC G9 Hom ospiens cDNA clone IMAGE:3080652 5' RGD-148089777 INIH MGC G9 Hom ospiens cDNA clone IMAGE:3080652 5' RGD-148089777 INIH MGC G9 Hom ospiens cDNA clone IMAGE:3080652 5' RGD-148089777 INIH MGC G9 Hom ospiens cDNA clone IMAGE:3080652 5' RGD-148089777 INIH MGC G9 Hom ospiens cDNA clone IMAGE:3080652 5' RGD-148089777 INIH MGC G9 Hom ospiens cDNA clone IMAGE:3080652 5' RGD-148080777 INIH MGC G9 Hom ospiens cDNA clone IMAGE:3080652 5' RGD-14808077 I
8979 18784 8979 18784 9008 19303 9176 18917		1.87	3.7E-01 XX 3.7E-01 UC 3.7E-01 QC	6677678 6677678 14982.1 1243525.1		Downe mayor for terminal decognicated to the property of the p
1 3	76	2.41		3.7E-01 D86976.1	MT	Human mRNA for KIAA0223 gene, partial cds

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-		_		_	_	_	_	_	_	_		_		_			_	_	_		_	_		_	_		_			_	_
	Top Hit Descriptor	DKFZp762K075_r1 762 (synonym: hmel2) Homo sapiens cDNA clane DKFZp762K075 5'	Homo sapiens NF2 gene	Bos taurus partial stat5B gene, exons 2-15 and Joined CDS	Human mibp gene, partial cds	yd03e05.r1 Soares infant brain 1NIB Homo sapleris cDNA clone IMAGE:24443 5'	yd03e05.r1 Soares Infant brain 1NIB Homo saplens cDNA clone IMAGE:24443 6'	hg33f02.x1 NCI_CGAP_GC6 Homo sapiens cDNA clone IMAGE:2947419 3'	hg33f02.x1 NCI_CGAP_GC6 Homo sapiens cDNA clone IMAGE:2947419 3'	Mus musculus ribosomal protein S19 (Rps19) gene, complete cds	Rattus norvegicus repeat element associated with the Rasgrf1 gene	Human mRNA for KIAA0323 gene, partial cds	P. Irregulare (P3804) gene for actin	RC5-ST0171-181099-011-g07 ST0171 Homo sapiens cDNA	PROTEIN LISOASPARTATE O-METHYLTRANSFERASE (PROTEIN BETA-ASPARTATE	METHYLIKANSFEKASE) (FIMI) (FKOTEIN C-SOASFAKTYL METHYLIKANSFEKASE) (F- ISOASPARTYL PROTEIN CARBOXYL METHYLTRANSFERASE)	Drosophila melanogaster sugar transporter 3 (sur3) mRNA, complete cds	H saplen's serotonin transporter gene, exons 9 and 10	H.sapiens serotonin transporter gene, exons 9 and 10	RC1-HT0545-150600-014-b12 HT0545 Homo sapiens cDNA	Brassica napus mRNA for MAP4K alpha2 protein	Bacteria from anoxic bulk soil 16S rRNA gene (strain XB45)	ha02g04.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:2872568 3'	Homo sapiens PHEX gene	yt74a06.r1 Soares fetal liver spleen 1NFLS Homo saplens cDNA clone IMAGE:275987 5'	wt72c10.x1 Soares_thymus_NHFTh Homo sapiens cDNA clone IMAGE:2513010 3' similar to TR:O15117 O15117 O15117 FYN BINDING PROTEIN. [1];	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 79	Homo sapiens lysosomal-associated membrane protein 2 (LAMP2), transcript variant LAMP2A, mRNA	Homo sapiens lysosomal-associated membrane protein 2 (LAMP2), transcript variant LAMP2A, mRNA	Homo sapiens chromosome 21 segment HS21C004	PROBABLE PEPTIDE ABC TRANSPORTER ATP-BINDING PROTEIN Y4TS
	Top Hit Database Source	EST_HUMAN	IN	TN.	M	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	TN	IN.	IN	IN	EST_HUMAN		SWISSPROT	FN	IN	IN	EST_HUMAN	NT	IN	EST_HUMAN	TN	EST_HUMAN	EST HUMAN	Z	TN	IN	TN	SWISSPROT
	Top Hit Acession No.	AL121154.1	Y18000.1	AJ237934.1	U89241.1	T80255.1	T80255.1	AW590184.1	AW590184.1	AF216207.1	AF056927.1	AB002321.1	X76725.1	AW812033.1		P24206	AF199485.1	X76758.1	X76758.1	BE707883.1	AJ009609.1	AJ229237.1	AW339393.1	Y10196.1	R94090.1	AW027174.1	AL161583.2	4504956 NT	4504956 NT		053194
	Most Similar (Top) Hit BLAST E Value	3.7E-01	3.7E-01	3.7E-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01		3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01		3.6E-01
	Expression Signal	2.18	1.52	1.31	71.7	2.59	2.59	6.05	9:05	6.18	1.08	1.04	2.39	1.83		2.28	8.9	1.85	1.85	1.16	0.85	1.23	2.01	1.6	5.49	1.68	13.42	3.37	3.37	1.4	15.34
	ORF SEQ ID NO:		25217				21055		21646	21682				22211		22350		23145	23146	23988	24312	24367		25928		26340		27269	27270		
	Exon SEQ ID NO:	19206	19251	19746	10902	11199	11199	11770	11770	11803	11899	12108	12221	12314		12460	15077	13341	13341	14205		14570	14819	15805	16104	16180	1	17082	17082	1_1	17524
	Probe SEQ ID NO:	9634	9701	9964	979	1292	1292	1874	1874	1908	2002	2223	2341	2437		2589	2869	3424	3424	4308	4835	4684	4941	2899	6238	6317	6816	7205	7205	7302	7674

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Top Hit Descriptor	601676418F1 NIH_MGC_21 Homo saplens cDNA clone IMAGE:3958997 5	Arabidopsis thaliana mRNA for SigB, complete cds	Methanobacterium thermoautotrophicum from bases 702375 to 714311 (section 62 of 148) of the complete	genome	Homo saplens hHb5 gene for hair keratin, exons 1 to 9	Escherichia coli K-12 MG1855 section 225 of 400 of the complete genome	Mus musculus Emr1 mRNA, complete cds	x60e11.x1 NCI_CGAP_Pen1 Homo sapiens cDNA clone IMAGE:2679116 3' similar to gb:K00658 TUBULIN ALPHA-1 CHAIN (HUMAN);	Mus musculus mannose receptor, C type 2 (Mrc2), mRNA	Arabidopsis thallana DNA chromosome 4, contig fragment No. 77	Homo sapiens GAP-like protein (LOC51306), mRNA	Homo sapiens GAP-like protein (LOC51306), mRNA	601811060R1 NIH_MGC_48 Home capiens cDNA clone IMAGE:4053951 3'	601894653F2 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:4124244 5'	Rattus norvegicus ADP-ribosylation factor-directed GTPase activating protein mRNA, complete cds	zr08a09.s1 Stratagene NT2 neuronal precursor 937230 Homo sepiens cDNA clone IMAGE:650872 3'	Fibrobacter succinogenes S86 endoglucanase E (celE) and endoglucanase D (celD) gene, complete cds	Danio rerio homeobox protein (hoxb5b) gene, complete cds	RC5-HT0218-181099-011-g02 HT0218 Homo sapiens cDNA	788iE1 fetal brain cDNA Homo saplens cDNA clone 788iE1-K similar to R07879, Z40498	Rat leukocyte common antigen (L-CA) gene, exons 1 through 5	S.scrofa mRNA for CD31 protein (PECAM-1)	Homo sapiens tumor protein p53-binding protein, 2 (TP53BP2), mRNA	Homo saplens tyrosine kinase non-receceptor 1 (TNK1), mRNA	VOLTAGE-DEPENDENT N-TYPE CALCIUM CHANNEL ALPHA-1B SUBUNIT (CALCIUM CHANNEL, L TYPE, ALPHA-1 POLYPEPTIDE ISOFORM 5) (ERAIN CALCIUM CHANNEL III) (BIII)	X.laevis gene for albumin including HP1 enhancer	C. griseus rhodopsin gene for opsin protein	Gallus gallus SPARC gene for osteonectin, promoter and exon 1	Gallus gallus SPARC gene for osteonectin, promoter and exon 1
Top Hit Database Source	EST_HUMAN	본		FN	ᅜ	Ā	NT	EST_HUMAN	ZI ZI	뒫	ᅜ	뒫	EST_HUMAN	EST_HUMAN	Ā	EST_HUMAN	Ā	N	EST_HUMAN	EST_HUMAN	NT	N	F	F	TORGSSIWS	LZ	Ę	L	NT
Top Hit Acession No.	BE902390.1	AB004293.1		AE000856.1	Y19210.1	AE000335.1	U66888.1	AW180229.1	TN 6678933 NT	AL161581.2	· 7706136 NT	7706136 NT	BF129796.1	BF310688.1	U35776.1	AA223252.1	U05897.1	AF071253.1	BE146585.1	N81203.1	M18349.1	X98505.1	11448042 NT	4507610 NT	002294				
Most Similar (Top) Hit BLAST E Value	3.6E-01	3.6E-01 A		3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.5E-01	3.5E-01	3.55-01	3.6E-01	3.5€-01	3.5E-01	3.5E-01	3.5E-01	3.5E-01	3.5€-01	3.5E-01	3.55-01	3.5€-01	3.6E-01	3.5年-01	3.5E-01	3 5E-01	3.5E-01	3.5E-01	3.5€-01	3.5E-01
Expression Signal	1.94	3.69		3.56	1.81	3.05	3.03	1.58	2.18	76.0	1.24	1.24	3.17	1.17	1.05	1.85	0.86	1.92	1.29	0.84	3.84	3.47	1.88	3.21	- 7.0	9	2.89	2.34	2.34
ORF SEQ ID NO:	28440			28134					19990							22327		23839						27238	276BD				
Exon SEQ ID NO:	18191	18326	L	17890	19761	18886	18983		10174	10598	L	L	10694	L	11526	12725	12537	14065	14275	L	14730		16584	L	17469	1	İ.,		1
Probe SEQ ID NO:	8314	8453		8741	9044	9127	9281	9889	202	662	708	708	763	1603	1622	2563	2672	4165	4379	4790	4849	6189	6704	7171	7640	7688	8116	8386	8386

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similer (Top) Hit BLAST E Vatue	Top Hit Acesslon No.	Top Hit Database Source	Top Hit Descriptor
8951	18758	29052	3.34	3.5E-01	AL161501.2	Ā	Arabidopsis thallana DNA chromosome 4, contig fragment No. 13
9208			2.12	3.5E-01	3.5E-01 X84565.1	NT	B.taurus atpA1 gene for F(0)F(1) ATP synthase alpha-subunit
9941	19616	24999		3.5E-01	H80814.1	EST_HUMAN	ys84f11.r1 Soares retina N2b4HR Homo sapiens ¢DNA clone IMAGE:218597 6'
9941	19816	25000		3.5E-01	3.5E-01 H80814.1	EST_HUMAN	ys64f11.r1 Soares retina N2b4HR Homo saplens cDNA clone IMAGE:219597 5'
691	10824		1.9	3.4E-01 AJ	AJ242956.1	<u> </u>	Homo saplens partal N-myc (exon 3), HPV45 L2 HPV45 L1, HPV45 E8, HPV45 E7 and HPV45 E1 genes isolated from IC4 central carcinoma cell line
98	10883	20731	4.53	3.4E-01	3.4E-01 Y09798.2	F	Pseudomonas fluorescens colR, colS genes, orf222 and partial inaA gene
1304	İ	21066		3.4E-01	3.4E-01 Y00554.1	Σ	Azotobacter vinelandii nifA gene for NifA protein (positive regulatory element)
2352	l _	22129		3.4E-01	3.4E-01 D90909.1	NT.	Synechocystis sp. PCC6803 complete genome, 1/1/27, 1311/235-1430418
2642	12509	22400	1.48	3.4E-01	3.4E-01 AL161516.2	NT	Arabidopsis thallana DNA chromosome 4, contig fragment No. 28
2971	1	22697	0.81	3.4E-01	AL163210.2	Ŋ	Homo saplens chromosome 21 segment HS21C010
2971	12898	22698	0.81	3.4E-01	3.4E-01 AL163210.2	NT	Homo saplens chromosome 21 segment HS21C010
3124	13049	22848	5.41	3.4E-01 U8	U83905.1	F	Canis familiaris rod photoreceptor cGMP-gated channel alphe-subunit (CNGC1) mRNA, complete cds
							Methylovorus sp. strain SS1 putative GrpE (grpE) DnaK (dnaK), and putative DnaJ (dnaJ) genes, complete
3488	13404	23209	3.48	3.4E-01 AF	AF106835.1	NT	cds
3722	13634		2.33	3.4E-01 BF	BF449010.1	EST_HUMAN	7n94e01.x1 NCI_CGAP_Ov18 Homo sapiens cDNA clone IMAGE:3572232 3' similar to TR:Q9UJ15 Q9UJ15 DJ18C9.1;
3965	13872		1.18	3.4E-01 AA	AA584196.1	EST_HUMAN	no11b10.s1 NCI_CGAP_Phe1 Homo sapiens cDNA clone IMAGE:1100347 3'
4394	14290	24074	0.78	3.4E-01	AF166341.1	NT	Homo saplens integrin alpha 6 (ITGA6) gene, exoits 12 through 23
4544	14437	24221	1.82	3.4E-01	3.4E-01 BE069912.1	EST_HUMAN	MR4-BT0403-230200-202-c01 BT0403 Homo sapiens cDNA
4828	14710	. 24494	0.93	3.4E-01	3.4E-01 BE463761.1	EST HUMAN	hy77d09x1 NCI_CGAP_GC6 Homo sapiens cDNA clone IMAGE:3197585 3' simitar to contains L1.t3 L1 repetitive alement;
	<u></u>						qi95c05.x1 NCI_CGAP_Kid3 Homo sapiens cDNA clone IMAGE:1867208 3' similar to contains Alu repetitive
4870	14750		3.57	3.4E-01 AI	AI240973.1	EST_HUMAN	element
4970	14845	24614	1.2	3.4E-01 X1	X16544.1	ħ	Sea urchlin hsp70 gene II for heat shock protein 70
5487	15406	25469	2.85	3.4E-01	AL161594.2	N	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 90
5551	1		4.71	3.4E-01	3.4E-01 AA085313.1	EST_HUMAN	zn12d11.s1 Stratagene hNT neuron (#937233) Homo saplens cDNA clone IMAGE:547221 3'
5654	15566		2.08	3.4E-01	3.4E-01 L02971.1	Ę	Echovirus 22 1AB, 1C, 1D, 2A, 2B, 2C, 3A, 3B, 3C, 3D proteins RNA, complete mature peptides and cds
5715		25724		3.4E-01	3.4E-01 AW204505.1	EST_HUMAN	UI-H-BI1-aei-e-12-0-UI.s1 NCI_CGAP_Sub3 Homo sapiens cDNA clone IMAGE:2719582 3'
5780	15687			3.4E-01	3.4E-01 AL120544.1	EST_HUMAN	DKFZp761A249_r1 761 (synonym: hamy2) Homo sapiens cDNA clone DKFZp761A249 5'
6045	ı		1.52	3.4E-01	N95225.1	EST_HUMAN	zb53e12.s1 Soares_fetal_lung_NbHL19W Homo capiens cDNA clone IMAGE:307342 3'

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
6148	16021	28161	1.41	3.4E-01	AI468082.1	EST_HUMAN	Im63g05.x1 NCI_CGAP_Brn25 Homo sapiens cDNA clone IMAGE:2162840 3' similar to gb:S37431 LAMININ RECEPTOR (HUMAN);
6943	16821		1.77		AA337063.1	EST_HUMAN	EST41765 Endometrial tumor Homo sapiens cDNA 5' end
7139	17016	27209	1.62	3.4E-01	9633624 NT	NT .	Bovine enterovirus strain K2577, complete genome
7313			3.88	3.4E-01	P26013	SWISSPROT	INTEGRIN BETA-8 PRECURSOR
7313	17189		3.88	3.4E-01	P26013	SWISSPROT	INTEGRIN BETA-8 PRECURSOR
7433			4.17	3.4E-01	U19482.1	IN	Saccharomyces cerevisiae Maf1p (WAF1) gene, complete cds
7433	16446	28636	4.17	3.4E-01	U19492.1	LN	Saccharomyces cerevisiae Maf1p (WAF1) gene, complete cds
7690	17540		2.22	3.4E-01	AJ225084.1	TN	Homo sepiens FAA gene, excn 16, 17 and 18
8380	18257		4.09	3.4E-01	AE000881.1	LΝ	Methenobacterium thermoautotrophicum from basis 1018444 to 1029212 (section 87 of 148) of the complete genome
8407	18283	28536	22	3.4E-01		SWISSPROT	PROBABLE E4 PROTEIN
8440	18314	28572	2.26	3.4E-01	AF045981.1	F	Rutilus arcasii cytochrome b (cytb) gene, mitochoridrial gene encoding mitochondrial protein, partial cds
8604	18471	28742	1.77	3.4E-01	M25856.1	NT	Human van Willebrand factor gene, exons 36 and 37
8604	18471	L	1.77	3.4E-01	M25856.1	N-	Human von Willebrand factor gene, exons 36 and 37
8790	18605	28895	1.8	3.4E-01	AB035507.1	ΝT	Rattus norvegicus mRNA for s-gicerin/MUC18, complete cds
8814	18627	28916	4.03	3.4E-01	AL161515.2	TN	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 27
9020	18814		2.01	3.4E-01	U93604.1	NT	Citrus variegation virus putative replicase gene, partial cds
9130	18887	,	1.3	3.4E-01	221621.1	N	S.cerevisiae RIB5 gene encoding Riboflavin synthase
9229	19508		1.82	3.4E-01	AF254351.1	TN	Schizosaccharomyces pombe Owf8p (cwf8) gene, complete cds
9347	19024		4.9	3.4E-01	126339.1	TN	Human autoantigen mRNA, complete cds
	ı						hw42h08.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3178127 3' similar to contains PTR5.f3
9378	-1		2.49	3.4E-01	BE21866	EST_HUMAN	PTK5 repetitive element;
9432			2.13			L	Beta vulgaris mitochondrion, complete genome
9537	19140	25264	1.79	3.4E-01	AJ297131.1	TN	Mus musculus SIL, MAP_17, CYP_a, SCL & CYP_b genes
							Homo saptens HLA class III region containing tenescin X (tenescin-X) gene, partial cds, cytochrome P460 21- hydroxylase (CVD24R), complement communent C4 (C4R) G41 helicase (SKI)W). RD, complement factor B
9829	19330		1.79	3.4E-01	AF019413.1	Z	(Bf), and complement component C2 (C2) genes,>
14	10000	19791	7.91	3.3E-01	X07990.1	LN	Rhizobium leguminosarum sym plasmid pRL5JI nodX gene
100	10000	19791	3.92	3.3E-01	X07890.1	LN	Rhizobium leguminosarum sym plasmid pRL5JI nodX gene
440	10384		1.16	3.3E-01	AL161545.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 45
617	ı		1.41	3.3E-01	7662485 NT	NT	Homo saplens KIAA1100 protein (KIAA1100), mRNA
1182	11093	20940	3.05	3.3E-01	Q12446	SWISSPROT	PROLINE-RICH PROTEIN LAS17

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Probe SEQ ID	Exan SEQ ID	ORF SEQ	Expression	Most Similar (Top) Hit	Top Hit Acession	Top Hit Database	Top Hit Descriptor
Ö	ö		Budio	Value	j Ž	Source	
1285	11193	21048	4.82	3.3E-01	BF568880.1	EST_HUMAN	602184016T1 NIH_MGC_42 Homo sapiens cDNA clone IMAGE:4300251 3'
1588	11492		1.11	3.3E-01	FN 58985 NT	¥	Mus musculus disintegrin 5 (Dtgn5), mRNA
1704			1.05	3.3E-01	AA332734.1	EST_HUMAN	EST36722 Embryo, 8 week I Homo saplens cDNA, 5' end
2355	12235		4.74	3.35-01	4507834 NT	1	Homo sepiens uridine monophosphate synthetase (orotate phosphoribosyl transferase and orotidine-5- decarboxylase) (UMPS) mRNA
2919	ı	22648	0		AJ25180	F	Bacteriophage phi-YeO3-12 complete genome
0000	i _		5	9 9E 01	000743	TORIGISIMS	INTERLEUKIN-12 ALPHA CHAIN PRECURSOR (IL-12A) (CYTOTOXIC LYMPHOCYTE MATURATION FACTOR 35 KD SUBUNIT) (CLMF P36)
30.48	_i_	75727	101		_	Ę	Streptomyces argillaceus mithramycin biosynthetic genes
3450	L	23173	1.14		_	Z	Homo sepiens MTA1-L1 gene, complete cds
3738	1		217	3.3E-01	084645	SWISSPROT	EXODEOXYRIBONUCLEASE V BETA CHAIN
3884	1		1.59	3.3E-01	AL161498.2	NT L	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 10
3920	L		1.92	3.3E-01		NT	Hypoxylon fragiforme chitin synthase gene, partial cds
4271	L		2.74	3.3E-01	D31662.1	NT	Rattus norvegicus DNA for regucalcin, partial cds
							to 78612.x1 NCL_CGAP_Ut3 Homo sapiens cDNA clone IMAGE:2205407 3' similar to gb.X57522 ANTIGEN
4589	14477		1.48			EST_HUMAN	PEPTIDE TRANSPORTER 1 (HUMAN);
5263	15185	24960	2.68			N	R.norvegicus mRNA for 3'UTR of ubiquitn-like protein
5263	15185	24961	2.68	3.3E-01		L	R.norvegicus mRNA for 3'UTR of ublqultin-like protein
5622	L		2.71	3.3E-01	BE618650.1	EST HUMAN	601472768T1 NIH_MGC_68 Homo saplens cDNA clone IMAGE:3875753 3'
5622	L		2.71	3.3E-01		EST_HUMAN	601472768T1 NIH_MGC_68 Homo sapiens cDNA clone IMAGE:3875753 3'
	ŀ				_		ty84h01.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2285809 3' similar to contains Alu
6117	16011	26148	3.64	3.3E-01	AI628131.1	EST HUMAN	repetitive element contains element. L'I repetitive element.
6117	16011	26149	3.64	3.3E-01	AI628131.1	EST_HUMAN	ty84h01.x1 NCI_CGAP_Kid11 Homo sapiens cDNA cione IMAGE:72285809 3' similar to contains Alu repetitive element;contains element L1 repetitive element;
8575	18433	26616	1.5	3.3E-01	N85146.1	EST HUMAN	J2498F Human fetal heart, Lambda ZAP Express Homo sapiens cDNA clone J2498 5' similar to TEGT
6984	ı		1			EST HUMAN	602140372F1 NIH_MGC_46 Home sapiens cDNA clone IMAGE:4301800 5'
7477	L	l			N69866.1	EST_HUMAN	za67h01.s1 Soares_fetal_lung_NbHL19W Homo sapiens cDNA clone IMAGE:297649 3'
7507	17295	l	2.81	3.3E-01	BF376745.1	EST_HUMAN	RC4-TN0077-250800-011-g04 TN0077 Homo saplens cDNA
7739			2.07	3.3E-01	L41044.1	NT	Homo sapiens high-mobility group phosphoprotein (HMGI-C) gene, exons 1-3, complete cds
8104	17994		2.71	3.3E-01	X63953.1	IN	D.maunttana Adh gene
8104	17994	28243	2.71	3.35-01	X63853.1	NT	D.mauritiana Adh gene
8389	18265		1.82		BF526499.1	EST_HUMAN	602070802F1 NCI_CGAP_Bm64 Homo sapiens cDNA clone IMAGE:4213585 5'
8561	18431	28700	12.76	3.3E-01	BE219351.1	EST_HUMAN	hv51g02.x1 NCI_CGAP_Lu24 Hamo sapiens cDNA clone IMAGE:3176978 3

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Table 4
Single Exon Probes Expressed in Heart

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acessian No.	Top Hit Database Source	Top Hit Descriptor
8662	18541	. 28826	я. 29.	3.3E-01	P47953	SWISSPROT	GALECTIN-3 (GALACTOSE-SPECIFIC LECTIN 3) (MAC-2 ANTIGEN) (IGE-BINDING PROTEIN) (35 KD LECTIN) (CARBOHYDRATE BINDING PROTEIN 35) (CBP 35) (LAMININ-BINDING PROTEIN) (LECTIN L-29) (CBP30)
8956	18763			3.3E-01	AA806621.1	П	ob71g02.s1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:1336850 3'
8972	10000	19791	1.88	3.3E-01	X07990.1	LN	Rhizobium leguminosarum sym plasmid pRL5JI nodX gene
9119	18881	28789	1.83	3.3E-01		TN	Homo sapiens aldehyde oxidase 1 (AOX1), mRNA
9817	19323		6.05	3.3E-01	AP000002.1	NT	Pyrococcus harikoshii OT3 genamic DNA, 28700(1-544000 nt. position (2/7)
449	10393		1.98	3.2至-01	AF018261.1	INT	Rattus norvegicus EH domain binding protein Epsin mRNA, complete cds
701	10834		2.05		í	TN	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 61
1148	11059	20902	18.89	3.2€-01	AF047013.1	LN	Fuserium poae virus 1 RNA2 putative RNA dependent RNA polymerase gene, complete cds
1261	11168	21019	1.04	3.2E-01	250202.1	IN	P.vulgaris arc5-1 gene
1368	11274	21130	6.25	3.2€-01	Q48624	SWISSPROT	LACTOSE PERMEASE (LACTOSE-PROTON SYMPORT) (LACTOSE TRANSPORT PROTEIN)
1737	11638	21506	1.22	3.2€-01	236041.1		S.cerevisiae chromosome II reading frame ORF YBR172c
1744	11645	21513	5.5	3.25-01	AW957194.1	EST_HUMAN	EST369284 MAGE resequences, MAGD Homo sapiens cDNA
1744	11645	21514	5.5		AW957194.1	EST_HUMAN	EST369264 MAGE resequences, MAGD Homo saplens cDNA
1800	11698	21574	1.16		AL111655.1	IN	Botrytis cinerea strain T4 cDNA library under conditions of nitrogen deprivation
2114	12003	21901	2.89	3.2E-01	BF203817.1	EST_HUMAN	601868804F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:4111512 5'
2484	12368		2.9	Ľ		TN	Mus musculus Pbx/knotted 1 homeobox (Pknox1); mRNA
2677	12542	22433			AF060568.1	N	Homo sapiens promyelocytic leukemia zinc finger protein (PLZF) gene, complete cds
3556	13470		96'0	3.2€-01	D10872.1	MT	Humam h NAT allele 3-2 gene for anylamine N-acétyltransferase
3868	13779		26.0	3.2€-01	AL161546.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 46
4295	14193	23977	1.35	3.2E-01	M18818.1	Ę	Rabbit beta-like globin gene cluster encoding the epsilon, gamma, delta (pseudogene) and beta globin polypeptides, complete cds
4375	14271	24052	0.81	3.2E-01	AF111167.2	NT.	Homo sapiens jun dimerization protein gene, partial cds; cfos gene, complete cds; and unknown gene
4397	14293	24077	1.33	3.2€-01	Q10268	SWISSPROT	HYPOTHETICAL 81.7 KD PROTEIN C13G7.04C IN CHROMOSOME I PRECURSOR
4632	14520		5.88	3.2E-01	BF693617.1	EST_HUMAN	602081972F1 NIH_MGC_81 Homo sapiens cDNA clone IMAGE:4246505 5
4807	15081		1.18	3.2E-01	M32352.1	IN	Mouse renin (Ren-1-d) gene, complete cds
5133	15000	24771	96.0		AY008847.1	Ы	Homo sapiens interleukin 12 p40 subunit (IL12B) gene, IL12B-1 allele, complete cds
5223	15146		282		BE173964.1	EST_HUMAN	CM0-HT0569-060300-269-f10 HT0569 Homo sapiens cDNA
6752	16631	26819			M60266.1	NT	Rat ISO-atrial natriuretic factor gene, complete cots
6872	16731	26946	13.32		X02508.1	Ā	H.sapiens gene fragment for acetylcholine receptor (AChR) alpha subunit exons 8, 9 and 3' flanking region
6875	16754	26951	14.29		_	EST_HUMAN	601897107F1 NIH_MGC_19 Homo saplens cDNA clone IMAGE:4126633 5

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Single Exon Probes Expressed in Heart

						200	
Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
6269	16807		1.35	3.2E-01	3.2E-01 AL161574.2	NT	Arabidopsis thallana DNA chromosome 4, contig fragment No. 70
9869	16863	27058	1.5	3.2E-01	3.2E-01 AE002015.1	IN	Delnococcus radiodurans R1 section 152 of 229 of the complete chromosome 1
7247	17124		2.19	3.2E-01	3.2E-01 M86511.1	NT	Human monocyte antigen CD14 (CD14) mRNA, complete cds
7741	17591	27812	ε	3.2E-01		NT	Borrelia burgdorferi plasmid cp32-2, erpC and erpD genes, complete cds, and unknown genes
7910	17760		3.25	3.2E-01	1,1	Ŋ	Homo sapiens gene for AF-6, complete cds
8028	17949	28199	2.7	3.2E-01	T06813.1	EST_HUMAN	EST04702 Fetal brain, Stratagene (cat#636206) Homo sapiens cDNA clone HFBDZ21
9152	19662		3.28	3.2E-01	L07288.1	Ę	Drosophila melanogaster laminin A (Lam-A) mRNA, complete cds
6996	19228		2:97	3.2E-01		SWISSPROT	ELONGATION FACTOR TU (EF-TU)
9801	18313		1.58	3.2E-01	39874.1	¥	Homo sapiens deoxycytidyiate deaminase gene, complete cds
9855	19698	24898	1.67	3.2E-01 B	BE385776.1	EST_HUMAN	60/1275480F1 NIH_MGC_20 Homo sapiens cDNA clone IMAGE:3616746 6'
							ye90h08.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:125051 5' similar to
2836	- 1	22397		3.1E-01	R18051.1	EST_HUMAN	gb:M64241 QM PROLEIN (HUMAN);
2665		22419		3.1E-01		۲	Homo sapiens KIAA0174 gene product (KIAA0174), mRNA
2665	12655	22420	3.45	3.1E-01	7661971 NT	NT	Homo sapiens KiAA0174 gene product (KIAA0174), mRNA
2827	12756		1.0.1		_	EST_HUMAN	hi46h08.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2975391 3'
3136			2.78		B022069.1	ΙN	Mus musculus gene for Ser/Thr kinase KKIAMRE, exon 6
3832	13744	23536		3.1E-01	AJ251586.1	TN	Daucus carota mRNA for transcription factor E2Fi(E2F gene)
5073	14943	24717	0.99	3.1E-01		EST_HUMAN	nm61h05.s1 NCI_CGAP_Br3 Homo sapiens cDNA clone IMAGE:10727613'
5385	15285	25119	8.88	3.1E-01		TN	Homo sapiens hepatocyte nuclear factor-3 alpha (HNF3A) gene, exon 1
5447	15368		45.03	3.1E-01		TN	Mus musculus mRNA for polycystin
5527		25510	1.94	3.1E-01	1	NT	Homo sapiens filamin 2 (FLN2) gene, exons 10 through 22
5890	15796	25918			3.1E-01 AW983549.1	EST_HUMAN	RC3-HN0001-310300-011-b04 HN0001 Homo sepiens cDNA
6160	19438	24861	2.4		3.1E-01 BE737392.1	EST_HUMAN	601306121F1 NIH_MGC_39 Homo sapiens cDNA clone IMAGE:3640420 5
7023		27092			R45318.1	EST_HUMAN	yg46f01.s1 Soares infant brain 1NIB Homo sapiens cDNA clone IMAGE:35639 3'
7791	L	27874			3.1E-01 BF69639.1	EST_HUMAN	602124743F1 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:4281811 5'
7791	17641	27875		3.1E-01 B	BF69639.1	EST_HUMAN	602124743F1 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:4281811 5'
. 7837	6824	8C0TC	80 1	2 40 04	3 15-03 01244001 1	NAMIN TOD	qi81e11.x1 NCI_CGAP_Kid3 Home sapiens cDNA clone IMAGE:1863980 3' similar to gb:S55700 HYDROXYMETHYI GI ITARYI -COA I YASE PRECI IRSOR (HI IMAN)
8211	L	2834B		1	BE218117 1	EST HIMAN	601883592F1 NIH MGC 57 Homo sapiens cDNA clone IMAGE:4095814 5
8823		28920	1 89		<u>!</u>	L	Homo sapiens KIAA0764 gene product (KIAA0764), mRNA
	l			1			Anolis opalinus Isolate QS NADH dehydrogenase subunit 2 (ND2) gene, complete cds; mitochondrial gene
9279	18982		1.63	3.1E-01	3.1E-01 AF294308.1	Ā	for mitochondrial product
9315	ii		1.64		3.1E-01 AF304162.1	NT	Stizostedion vitreum 40S ribosomal protein S11 mRNA, partial ods
9457	19089		2.45		AF195953.1	·	Homo sapiens membrane-bound aminopeptidase P (XNPEP2) gene, complete cds

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Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Vatue	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
000						ţ	Homo sapiens transcription factor IGHM enhancer 3, JM11 protein, JM4 protein, JM5 protein, T54 protein, JM10 protein, A4 differentiation-dependent protein, triple LIM domain protein 6, and synaptophysin genes,
9080	Į	İ			-196//9.1	Z	complete cos, and L-type calcium channel a>
8					6765083 NT	뒫	Mus musculus protein kinase C, epsilon (Pkce), mRNA
264			11.12			F	Homo sapiens Xq pseudoautosomal region; segment 1/2
1204	11114				3.0E-01 AW300400.1	EST_HUMAN	xs63f08.x1 NCI_CGAP_Kld11 Homo saplens cDNA clone IMAGE:2774343 3'
1491	11396	21256			1	TN.	Balaenoptera physalus gene encoding atrial natriuretic peptide
2089	11978	21873				Ę	Rattus norvegicus Ca2+/calmodulin-dependent protein kinase II, alpha subunit mRNA, 3' untranslated region
3175	13100					F	Corynebacterium sp. ALY-1 alyPG gene for polyguluronate lyase, complete cds
3789	13701	23488	1.34	L	1	EST_HUMAN	PM1-ST0262-261189-001-g01 ST0262 Homo sapiens cDNA
4412	14306	24089		L	3.0E-01 AJ006755.1	IN	Balaenoptera physalus gene encoding atrial natriuretic peptide
5283	15205	24981	5.34	L		EST_HUMAN	601594960F1 NIH_MGC_9 Homo septens cDNA clone IMAGE:3948734 5'
5379	,					EST_HUMAN	RC3-BT0333-180700-111-a03 BT0333 Homo saplens cDNA
5379						EST_HUMAN	RC3-BT0333-180700-111-a03 BT0333 Homo sapiens cDNA
5401						IN	Mus musculus 129/sv Clara cell 10 kd protein (mCC10) gene, complete cds
9809					D16313	M	Mouse cytokeratin 15 gene, complete cds
6335	16198	26358	2.57	3.0E-01	10947007 NT	NT.	Mus musculus midnolin (Midn-pending), mRNA
6428	16290	26451	1.35		3.0E-01 AF071810.1	TN	Streptococcus pneumoniae strain DBL5 PspA (pspA) gene, partial cds
6625	16506	26693	1.25		3.0E-01 AE001755.1	IN	Thermotoga maritima section 67 of 136 of the complete genome
2077	03504		107	9000	TIA	Ŀ	Mus musculus C-type (calcium dependent, carbohydrate recognition domain) lectin, superfamily member 9
6837	L	27007			BF566083 1	EST HUMAN	601339079F1 NIH MGC 53 Hamo sepiens cDNA clone IMAGE:3681594 5
7944	<u>_</u>		1.89			E	Aspergillus oryzae bipA gene for ER chaperone BiP, complete cds
8980	18785				3.0E-01 H51029.1	EST_HUMAN	yp84b10,r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:194107 5'
8880	18785	28075	2.89		3.0E-01 H51029.1	EST_HUMAN	yp84b10.r1 Soares fetal liver spleen 1NFLS Home sapiens cDNA clone IMAGE:194107 5'
9564	19647		1.43		3.0E-01 AJ297631.1	NT	Rettus norvegicus mRNA for glyceraldehyde-3-phosphate dehydrogenase type 2 (gapdh-2 gene)
9836	.19690		2.76		TN 6677766 NT	NT.	Mus musculus ribose 5-phosphate isomerase A (Rpia), mRNA
1978	11871				2.9E-01 AE000736.1	NT	Aquifex ecolicus section 68 of 109 of the complete genome
2201		21989				ᅜ	Chrysodidymus synuroideus mitochondrion, complete genome
3147						F	Xenopus laevis transcription factor E2F mRNA, complete cds
3213					2.9E-01 AW754239.1	EST_HUMAN	PM1-CT0326-171299-001-f12 CT0326 Homo sapiens cDNA
3213	13137	22940	2.29		2.9E-01 AW754239.1	EST_HUMAN	PM1-CT0328-171299-001-112 CT0326 Homo sapiens cDNA

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Top Hit Database Source	TOTAT1 X1 NCI_CGAP_Gas4 Homo sapiens cDNA clone IMAGE:2188412 3' similar to gb:D15050 NIL-2-A EST_HUMAN ZINC FINGER PROTEIN (HUMAN);contains element L1 repetitive element;	EST_HUMAN wr02r10.x1 NC _CGAP_GC6 Homo sapiens cDi\A clone IMAGE:2480395 3	П	EST HUMAN repetitive element	NT Mus musculus SKD1 (Skd1) gene, complete cds	NT (Mus musculus SKD1 (Skd1) gene, complete cds	NT Mus musculus gene, complete cds, similar to EXLM1	EST_HUMAN MT7e12.s1 Soares Infant brain 1NIB Homo sapiehs cDNA clone IMAGE:28291.3'		NT system polypeptides P16,18,28,30 and levanase;	B.subtilis levanase operon levC, levE, levE, levC and sacC (partial) genes for fructose phosphotransferase system polypeptides P16.18.28.30 and levanase.			NT [Mus musculus Filih protein (Filih) gene, complete cds; and Light protein (Light) gene, partial cds	ISSPROT	Mus musculus major histocompatibility locus class II region: Fas-binding protein Daxx (DAXX) gene, partial code. Blind (BING4) Blind (BING4) heizh 3.	NT galactosyl transferase (beta1,3-galactosyl tr>	EST_HUMAN 601065830F1 NIH_MGC_10 Homo sepiens cDNA clone IMAGE:3462287 6'	EST_HUMAN 801085830F1 NIH_MGC_10 Home sapiens cDNA clone IMAGE:3452287 5'	NT Trypanosoma cruzi stage-specific surface glycoprotein gp82 (gp82) mRNA, partial cds	NT Torpedo californioa mRNA enooding acetylcholine receptor gamma subunit	NT Torpedo californica mRNA encoding acety/choline receptor gamma subunit	InyS5h02.s1 NCI_CGAP_Pr12 Homo sapiens cDNA done IMAGE:1273779 similar to contains LTR8.t2 LTR8	NT Campylobacter jejuni NCTC11168 complete genome; segment 5/6	1	T_HUMAN	NT Homo sapiens TNF-a-inducible RNA binding protein (TIRP) gene, complete cds	T_HUMAN	NT Chlamydomonas reinhardtii mRNA for nitrite reductase structural locus	NT Chlamdomonas reinhardtii mRNA for nitrile reductase structural locus
Top I Dateb	EST_HUN	EST_HUN		EST HUN	Ę	Έ	Ę	EST_HU	!	Z	5		Ę	Ę	SWISSPF		Þ	EST_HUN	EST_HUN	卜	Ā	NT	110 100			EST_HUN	된	EST_HUM	N.	F
Top Hit Acession No.	2.9E-01 AI610836.1	2.9E-01 AW002902.1			П	2.9E-01 AF134119.1		2.9E-01 R37485.1		2.9E-01 X56098.1	2 SE-01 X56098 1	662	2.9E-01 U03420.1	Ī						F.		2.9E-01 V01394.1		2.9E-01 At 139078.2				3.1		2 0E_01 V08037 1
Most Similar (Top) Hit BLAST E Value	2.9E-01	2.9E-01		2.8E-01	2.9E-01	2.9E-01	2.9E-01	2.9E-01	1	2.9E-01	2 SF-01	2.9E-01	2.9E-01	2.9E-01	2.9E-01		2.9E-01	2.9E-01	2.9E-01	2.9E-01	2.9E-01	2.9E-01	20 00	2.8E-01		2.9E-01	2.9E-01	2.9E-01	2.9E-01	2 9F-01
Expression Signal	1.28	0.82		1.01	0.78	0.78	0.92	1.49		4.46	4 45	6.83	2.24	1.51	2.52		1.74	1.76	1.78	1.96	2.69	2.69	,	454		1.63	2.74	1.33	4.86	4 88
ORF SEQ ID NO:	23522			24059	24064	24065	24401			25504	25505	25513	25838	24847	26234		26267	26686	26687	28387	28639	28640	39090	28967		25260	25248		25188	25189
SEQ ID	13733	13901		14280	14284	14284	14615	15141		15440	15440	15448	15724	15128	16084		16115	16499	16499	18147	18375	18375	40678	18878		19127	19184	19212	19367	19387
Probe SEQ ID NO:	3821	3894		4384	4388	4388	4729	5218		5522	5522	6529	5818	6171	6218		6249	6619	6819	8267	8502	8502	7300	8888		9514	9802	9644	9883	9883

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Single Exon Probes Expressed in Heart

Top Hit Descriptor	Rattus norvegicus A-kinase anchoring protein AKAP150 mRNA, complete cds	Guira guira oocyte maturation factor Mos (c-mos) gene, partial cds	601148733F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:3163688 5	601148733F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:3163688 5'	Human mRNA for serine/threonine protein kinase, complete cds	QV1-CT0364-120200-065-b05 CT0364 Homo sapiens cDNA	DKFZp586I2321_r1 586 (synonym: hute1) Homo sapiens cDNA clone DKFZp586I2321	hd44b03.x1 Soares_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:2912333 3'	Escherichia coli K-12 MG1655 section 384 of 400 of the complete genome	Escherichia coli K-12 MG1655 section 384 of 400 of the complete genome	Arabidopsis thaliana DNA chromosome 4, contigi fragment No. 65	Arabidopsis thaliana mRNA for lipoyftransferase, complete cds	Toxoplasma gondii 90kDa heat-shock protein (HSP90) mRNA, partial cds	B.taurus microsatellite (ETH121)	B.taurus microsatellite (ETH121)	Pyrococcus harikoshii OT3 genomic DNA, 777001-894000 nt. positian (4/7)	Borrelia burgdorferi (section 66 of 70) of the complete genome	ov44g10.x1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1640226 3' similar to contains Alu repetitive element contains element MER22 repetitive element;	Mus musculus chromosome X contigA; putative Magea9 gene, Caltractin, NAD(P) steroid dehydrogenase	and Zinc finger protein 185	RNA POLYMERASE BETA SUBUNIT (LARGE STRUCTURAL PROTEIN) (L PROTEIN)	Human mRNA for transcription factor AREB6, complete cds	Human mRNA for transcription factor AREB6, complete cds	Bowne adenowirus 3 complete genome	602042601F1 NCI_CGAP_Brn67 Homo sapiens,cDNA clone IMAGE:4180129 6'	q159c11.x1 Soares_NhHMPu_S1 Homo sapiens cDNA clone IMAGE:1876628 3' similar to contains Alu	repetitive element; contains element LTR5 repetitive element;	EST57072 Infant brain Homo saplens cDNA 5' end	Homo sapiens OCTN2 gens, complete cds	Mesembryanthemum crystallinum fructose-biphosphate aldolase mRNA, complete cds	Mesembryanthemum crystallinum fructose-biphosphate aldolase mRNA, complete cds	UI-H-BI4-aci-f-04-0-UI.s1 NCI_CGAP_Sub8 Horrio sapiens cDNA clone IMAGE:3085182 3'
Top Hit Database Source	NT TA	TN.	EST_HUMAN	EST_HUMAN	ᄓ	EST_HUMAN	EST_HUMAN	EST_HUMAN	TN	TN	Z	LN LN	N	N	NT	N	Z	EST HUMAN		뉟	SWISSPROT	TN	TN	TN	EST_HUMAN		EST_HUMAN	EST_HUMAN	TN	TN	Z	EST_HUMAN
Top Hit Acession No.	U67136.1	AF168050.1	BE313442.1	BE313442.1	D86550.1	AW860020.1	AL047620.1	AW511195.1	AE000494.1	AE000494.1	AL161565.2	AB020975.1	AF179480.1	Z14037.1	Z14037.1	AP000004.1	AE001180.1	A1090868.1		AL021127.2		D15050.1		AF030154.1	BF528188.1		AI272669.1	AA348997.1	AB016625.1	AF003124.1	AF003124.1	BF511215.1
Most Similar (Top) Hit BLAST E Value	2.8E-01	2.8E-01	2.8E-01	2.8E-01	2.8E-01	2.8E-01	2.8E-01	2.8E-01	2.8E-01	2.8E-01	2.8E-01	2.8E-01	2.8E-01	2.8E-01	2.8E-01	2.8E-01	2.8E-01	2.8E-01		2.8E-01	2.8E-01	2.8E-01	2.8E-01	2.8E-01	2.8E-01		2.8E-01	2.8E-01	2.8E-01	2.8E-01	2.8E-01	2.8E-01
Expression Signal	1.84	2.47	1.1	1.1	1.01	1.55	1.77	1.47	2.94	2.94	2.47	1.37	1.49	2.48	2.48	1.1	1.75	2.21		0.98	22	1.03	1.03	2.8	45.L		1.7	21.38	2.33	1.44	1.44	7.67
ORF SEQ ID NO:		20828	21013	21014	21025	21470	21750	21869		22198		22388		22667	89977	23057	23604			24034	24038			24394	24427		24459	24948	25419	25784		26073
Exon SEQ ID NO:	10497	10983	11163	11163	11176	11598	11860	11974	12300	12300	12375	12498	12867	12868	12868	13252	13824	14003		14248	14253	14559	14559	14608	14640			19440	15363			15941
Probe SEQ ID NO:	556	1067	1258	1256	1269	1698	1968	2084	2423	2423	2500	2630	2940	2941	2941	3332	3915	4103		4352	4357	4673	4673	4722	4755		4787	6262	5443	92.20	6770	6038

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
6713	16593	26781	1.24	2.8E-01	Al346126.1	EST_HUMAN	qp48h01x1 NCI_CGAP_Co8 Homo sepiens cDNA clone IMAGE:1926289 3' aimilar to gb:X06323_cds1 MITOCHONDRIAL 60S RIBOSOMAL PROTEIN L3 (HUMAN);
6713	16593	26782	1.24	2.8E-01	Al346126.1	EST_HUMAN	qp48h01.x1 NCI_CGAP_Co8 Homo sapiens cD\NA clone IMAGE:1926289 3' similar to gb:X08323_cds1 MITOCHONDRIAL 60S RIBOSOMAL PROTEIN L3 (HUMAN);
8778	16657	26847	2.39	2.8E-01	U51688.1	TN	Homo sapiens lanosterol 14-alpha demethylase cytochrome P450 (CYP51) gene, exon 5
6994	16871		7.25	2.8E-01	BF347847.1	EST_HUMAN	602022987F1 NCI_CGAP_Bm67 Homo saplens cDNA clone IMAGE:4158528 5'
7311	17187	27388	1.35	2.8E-01	AF080592.1	NT	Mus musculus centin (Cetn2) gene, complete cds
7600	17451		1.15	2.8E-01	L13654.1	NT	Lycopersicon esculentum peroxidase (TPX1) mRNA, complete cds
7789	17639	27872	2.79	2.8E-01		NT	Homo sapiens hypothetical protein (LOC\$1319), mRNA
8126	18014	28261	2.27	2.8E-01	BF241062.1	EST_HUMAN	601880794F1 NIH_MGC_55 Homo sapiens cDNA clone IMAGE:4109350 5'
8126	18014	28262	2.27	2.8E-01	BF241062.1	EST_HUMAN	601880794F1 NIH_MGC_55 Homo sapiens cDNA clone IMAGE:4109350 5'
8153	18041	28291	3.01	2.8E-01	-	EST_HUMAN	801852148F1 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:4076026 5'
							Drosophila heteroneura fruitless (fru) gene, alternative splice products, 5' flanking region, exons 1 through 7
8254	18134	28382	2.62	2.8E-01	AF051662.1	NT	and complete cds
8238	18466		3.49	2.8E-01	BF674023.1	EST_HUMAN	602137418F1 NIH_MGC_83 Homo sapiens cDNA clone IMAGE:4273853 5'
9552	19150		26.9	2.8E-01	D83329.1	IN	Mus musculus DNA for prostaglandin D2 synthase, complete cds
9657	19219	25236	3.11	2.8E-01	BE178699.1	EST_HUMAN	PM4-HT0606-030400-001-a07 HT0606 Homo sapiens cDNA
9820	19851		1.69	2.8E-01	11433629 NT	TN	Homo saplens CDC42-binding protein kinase beta (DMPK-like) (CDC42BPB), mRNA
468	10411	20230	2.33	2.7E-01	Y17324.1	LN	Rattus norvegicus CDK104 mRNA
							z39b10.s1 Soares_total_fetus_Nb2HF8_9w Horno sapiens cDNA clone IMAGE:788827 3' similar to
597	10533		3.14	2.7E-01	~	EST_HUMAN	contains Alu repetitive element;
1240	11147	20896	1.49	2.7E-01	AB004906.1	NT	lpomoea purpurea transposable element Tip100 gene for transposase, complete cds
1604	11509		1.73	2.7E-01	X79815.1	LN	G.lamblia SR2 gene
1698	11600	21471	2.96	2.7E-01	W 58067.1	EST_HUMAN	zd22h10.r1 Soares_fetal_heart_NbHH19W Homo sapiens oDNA clone IMAGE:341443 6'
							GAG POLYPROTEIN (CONTAINS: INNER COAT PROTEIN P12; CORE PROTEIN P15; CORE SHELL
1738		21507	1.25	2.7E-01	P03341	SWISSPROT	PROTEIN P30; NUCLEOPROTEIN P10]
2088	12712		2.38	2.7E-01	AF047575.1	L	Rattus norvegicus vesicular monoamine transporter type 2, promoter region and exon 1
2316	12197	22094		2.7E-01	Y13868.1	INT	Feline immunodeficiency virus env gene, isolate ITTO088PIU (M88), partial
							ta43c11.x2 NCI_CGAP_Lu25 Homo sapiens cDNA clone IMAGE:2046836 3' similar to contains element L1
2408	12283	22180	3.51	2.7E-01	AI310858.1	EST_HUMAN	repetitive element;
2856			1.25	2.7E-01	BF088284.1	EST_HUMAN	CM1-HT0875-060900-385-e05 HT0875 Homo sapiens cDNA
3929				2.7E-01	Al928015.1	EST_HUMAN	wo92e11.x1 NCI_CGAP_Kld11 Homo saplens cDNA clone IMAGE:2462828 3'
3943				2.7E-01	L 77 569.1	L	Homo sapiens DiGeorge syndrome critical region ∣telomeric end
4798	14681	24468	0.89	2.7E-01	_27516.1	NT	Triticum aestivum (Wcs66) gene, complete cds

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Probe	Exon	CHA	Evareacion	Most Similar	Top Hit Acession	Top Hit	Top Hit
SEQ ID	SEQ ID	ον σ ON O:	Signal	BLAST E	No.	Database Source	Top Hit Descriptor
4973	14848		3.5	2.7E-01	AW856131.1	EST_HUMAN	RC1-CT0286-230200-016-e03 CT0286 Homo sapiens cDNA
5221	15144	. 24838	3.49	2.7E-01	P17277	SWISSPROT	HOMEOBOX PROTEIN HOX-A4 (CHOX-1.4)
6064	16047	26192	225	2.7E-01	Q61554	SWISSPROT	FIBRILLIN 1 PRECURSOR
6527	16386	26565	2.23	2.7E-01	AF248054.1	F	Bos taurus micromolar calcium activated neutral protease 1 (CAPN1) gene, exons 11-20, and partial cds
6527	16386	28566	2.23	2.7E-01	AF248054.1	Ę	Bos taurus micromolar calcium activated neutral protease 1 (CAPN1) gene, exons 11-20, and partial cds
7378	17247	27463	10.08	2.7E-01		SWISSPROT	THREONYL-TRNA SYNTHETASE (THREONINE-TRNA LIGASE) (THRRS)
7378	ļ		10.08	2.7E-01	_	SWISSPROT	THREONYL-TRNA SYNTHETASE (THREONINETRNA LIGASE) (THRRS)
7379	ļ		2.24	2.7E-01	P37928	SWISSPROT	FIMBRIAE W PROTEIN
7801	17651	27888	1.29	2.7E-01	AF091848.1	M	Oryctolagus cuniculus calgranulin C mRNA, partial cds
7827	17677	27921	1.93	2.7E-01	AF087434.1	TN	Mus musculus transcription factor NF-ATc Isoform a (NF-ATca) mRNA, complete cds
8187	18073	28322	1.76	2.7E-01	AV705043.1	EST_HUMAN	AV705043 ADB Homo sapiens cDNA clone ADBCOD05 5'
8187	18073	28323	1.76	2.7E-01	AV705043.1	EST_HUMAN	AV705043 ADB Homo sapiens cDNA clone ADBCOD05 6'
1070	70007	20200	97	27 75 0	A 1422280 4	LV.	Homo sapiens caveolin-1/-2 locus, Contig1, D7S522, genes CAV2 (exons 1, 2a, and 2b), CAV1 (exons 1 and
9368	L.	20002	1.27	2.7E-01	X95267.1	, LZ	G.galius mRNA for ryanodine receptor type 3
9810	1		2.15	2.7E-01	AF217491.1	Ā	Homo sapiens fragile 16D oxido reductase (FOR) gene, exon 6
462	l	20224	1.54	2.6E-01	P78411	SWISSPROT	IROQUOIS-CLASS HOMEODOMAIN PROTEIN IRX-2
472	10418		1.39	2.6E-01	D16459.1	Ę	Bos taurus mRNA for mb-1, complete cds
1371	11277	21133	2.23	2.6E-01	BE885087.1	EST_HUMAN	601510838F1 NIH_MGC_71 Homo sepiens cDNA clone IMAGE:3912345 5
1418	11324	21188	76.0	2.6E-01	AB013290.1	NT	Glycine max pseudogene for Bd 30K
1854	11750	21624	8.69	2.6E-01		NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 2
1854	11750	21625	8.59	2.6E-01	AL161472.2	TN	Arebidopsis thallana DNA chromosome 4, contig fragment No. 2
							bb04d10.x1 NIH_MGC_14 Homo sapiens cDNA done IMAGE:2958451 3' similar to gb:M36072 60S RIBOSOMAL PROTEIN L7A (HUMAN); gb:M14689_cds1 Mouse surfeit locus surfeit 3 protein gene
2046	11937		6.6	2.6E-01	AW 733152.1	EST_HUMAN	(MOUSE);
2106	11995	21895	1.5	2.6E-01	M11844.1	F	Human prealburnin gene, complete cds
2424	12301		2.68	2.6E-01	Y12996.1	۲	B.martitmus rbcL gene
2499	L		9.68	2.6E-01	BE272440.1	EST_HUMAN	601126016F1 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:2990043 5
3053	12980		1.03	2.6E-01	AW974531.1	EST_HUMAN	EST386635 MAGE resequences, MAGM Homo éapiens cONA
3485	13401		0,94	2.6E-01	BE217816.1	EST_HUMAN	hv30c02.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3174914 3' similar to contains L1.t3 L1 repetitive element;
3532	ł	23245		2.6E-01	M22342.1	N	Bacteriphage T2 DNA-(adenine-N6)methyltransferase (dam) gene, complete cds

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					5	30 - 100 - 10F	July Charles Lypressed III realt
Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Toʻɔ Hit Descriptor
3591	13505	23295	1.7	2.6E-01	AF229118.1	LN	Homo sapiens acetylcholinesterase collagen-like tail subunit (COLQ) gene, exons 1A, 2, 3, 4, and 5
3651	13565	23351	6.0	2.6E-01	4B017446.1	NT	Rattus norvegicus mRNA for organic anion transporter 3, complete cds
4010	13916		+	2.6E-01	4W859510.1	EST_HUMAN	EST371580 MAGE resequences, MAGF Homo sapiens cDNA
4062	13984		13.13	2.6E-01	BE080598.1	EST_HUMAN	QV1-BT0630-040400-132-e03 BT0630 Homo sapiens cDNA
į	1		.00	20.0	, 00000		Enterococcus faecium strain N97-330 vanD glycopeptide resistance gene cluster, complete cds; and
4239	14158		0.85	2.6E-01	AF175293.1	IN.	unknown gene C-II. o z-II. o z-Divi (c- z-II. d-II. z- z- z- z- z- z- z- z- z- z- z- z- z-
4393	14289		0.84	2.6E-01	AB021180.1	NT	Gallus gallus mKNA tor skeletal myosin neavy chain, complete cds
. 4393	14289		0.84	2.6E-01	AB021180.1	NT	Gailus gallus mRNA for skeletal myosin heavy chain, complete cds
4448	14342	24134	1.17	2.6E-01	AA457617.1	EST_HUMAN	aa89d07.r1 Stratagene fetal retina 937202 Homo sapiens cDNA clone IMAGE:838477 5'
4548	14441	24224	1.63	2.6E-01	U01103.1	NT	Arabidopsis thaliana PSI type III chlorophyll a/b-binding protein (Lhca3*1) mRNA, complete cds
4616	14504	68676	1 28	2 BE-01	AF142703 1	TV.	Obbresita radicosa maturase-like protain (matX) dene, complete cds; chloroplast gene for chloroplast product
7808	14778	L	2 56	2 BE.04	HOARER 1	EST HIMAN	451e05 rt Scares placenta Nb2HP Homo seplens cDNA clone IMAGE:152288 5
5051	14923		0.89	2.6E-01	-	SWISSPROT	ACYL-COA DEHYDROGENASE, MEDIUM-CHAIN SPECIFIC, MITOCHONDRIAL PRECURSOR (MCAD)
002.5	10761	L	60.0	2000	AE004044 4	TIA	Thermotone modifine continu 192 of 138 of the rivinglate nearons
3	19/04		2.03	Z.0E-U1	AEUU1811.1	Z	The mooga manning section 125 of the complete genotice
5763	15670	25777	1.93	2.6E-01	AI582557.1	EST_HUMAN	ts02e12.x1 NCI_CGAP_Pan1 Homo sapiens cDNA clone IMAGE:2227438 3' similar to SW:NDF1_RAT Q84289 NEUROGENIC DIFFERENTIATION FACTOR 1 ;contains element LTR1 repetitive element ;
	ļ		,	i i			ts02e12.x1 NCI_CGAP_Pan1 Homo saplens cDNA clone IMAGE:2227438 3' similar to SW:NDF1_RAT
20/62	130/0	81/67	1.93	Z.0E-U1	Alb62557.1	EST TOWAR	487263 VECNOCERNO DIFFERENTIALINY FACTOR 1 JUNIORIEM SEGMENT ET N. LEPENDA SEGMENT.
6552	16410	26588	1.52	2.6E-01	R10365.1	EST_HUMAN	gb:X12517 U1 SMALL NUCLEAR RIBONUCLEÓPROTEIN C (HUMAN);
6583	16465	26658	1.27	2.6E-01	R02411.1	EST_HUMAN	ye82a07.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:1242125'
6852	16731	26925	3.01	2.6E-01	BF343588.1	EST_HUMAN	602014422F1 NCI_CGAP_Bm64 Homo sapiens cDNA clone IMAGE:4150396 5'
0069	16779		2.04	2.6E-01	Q10199	SWISSPROT	HYPOTHETICAL 75.2 KD PROTEIN C11C11.02 IN CHROMOSOME II
7046	16923	27112	4.34	2.6E-01	BE830339.1	EST_HUMAN	RC5-ET0082-310500-021-F10 ET0082 Homo sapiens cDNA
7046	16923		4.34	2.6E-01	BE830339.1	EST_HUMAN	RC5-ET0082-310500-021-F10 ET0082 Homo sapiens cDNA
7854	17704		1.16	2.6E-01	0.28295	SWISSPROT	VON WILLEBRAND FACTOR PRECURSOR (VWF)
8812	18625		93.65	2.6E-01	X51755	NT	Human lambda-immunoglobulin constant region complex (germline)
9136	18892		1.98	2.6E-01	10190655	LN	Mus musculus jerky (Jrk), mRNA
9328	1		1.92	2.6E-01	3E883491.1	EST_HUMAN	601511052F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3912612 5
9395	19053	25309	2.6	2.6E-01	AF316896.1	Ę	Homo sapiens Na/K-ATPase gamma subunit (FX/D2) gene, complete cds, alternatively spliced

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Probe SEQ ID NO: 9833 9889 9889 1044 1105 1105 1105 1105 1242 242 242 242 244 1697 1697 1697 1746 2367 246 2367 246 2367 246 2367 246 246 246 246 246 246 246 246		ORF SEQ ID NO: 20025 20026 200	Expression Signal 1.5 6.03 6.03 6.03 6.03 6.03 6.03 6.08 6.09 6.85 6.09 6.86 6.09 6.86 6.09 6.86 6.09 6.88 6.29 7.2.87 7.83 6.08 6.08 6.08 6.09 6.09 6.09 6.09 6.09 6.09 6.09 6.09	Most Similar (Top) Hit BLAST E Vabue 2.6E-01 2.6E-01 2.5E-01 2		Top Hit Database Source Source Source Source NT NT NT NT NT NT NT NT NT NT NT NT NT	Homo sapiens inositol polyphosphale 1-phosphalase (INPP1) gene, complete cds ADENYLATE CYCLASE (ATP PYROPHOSPHATE-LYASE) (ADENYLAT CYCLASE) ADENYLATE CYCLASE (ATP PYROPHOSPHATE-LYASE) (ADENYLAT CYCLASE) gene encoding mitochordial protein, mRNA Homo sapiens ATP synthase, H+ transporting, mitochondrial F1 complex, delta subunit (ATP5D), nuclear gene encoding mitochordial protein, mRNA Homo sapiens ATP synthase, H+ transporting, mitochondrial F1 complex, delta subunit (ATP5D), nuclear gene encoding mitochordrial protein, mRNA Starfish (P. cohraceus) cytoplasmic actin gene, complete cds Was musculus (GR/Swiss glyceradehyde 3-phosphate dehydrogenase (Gapd-S) gene, complete cds Ureaplasma urealydrum section 57 of 59 of the complete genome yert gg/7 x1 Stratagene lung (#837210) Homo sapiens cDNA clone IMAGE:117486 of Bohydis cherea strain 74 cDNA library under conditions of nitrogen deprivation Homo sapiens hyperpolarization activated cyclic nucleotide-gated potasstum channel 4 (HCN4) mRNA PMA-CT0400-310700-005-408 CT0400 Homo sapiens cDNA Aquitex eeolicus section 7 of 109 of the complete/genens cDNA Aquitex eeolicus section 7 of 109 of the complete/genens cDNA Aquitex eeolicus section 7 of 109 of the complete/genens cDNA EST385464 MAGE resequences, MGM Homo sapiens cDNA Blaurus mRNA for D-aspartale oddsee EST385444 MAGE resequences, MGM Homo sapiens cDNA Aradidopsis thaliana DNA chromosome 4, contig fragment No. 29 wg11c07 x1 Socree_NSF F8 gW_OT PA_PS; Homo sapiens cDNA clone IMAGE:2364780 3' wg11c07 x1 Socree_NSF F8 gW_OT PA_PS; Homo sapiens cDNA clone IMAGE:2364780 3' Wg11c07 x1 Socree_NSF F8 gW_OT PA_PS; Homo sapiens cDNA clone IMAGE:2364780 3' Wg11c07 x1 Socree_NSF F8 gW_OT PA_PS; Homo sapiens cDNA clone IMAGE:2364780 3' Wg11c07 x1 Socree_NSF F8 gW_OT PA_PS; Homo sapiens complete cds; and Naip3 gene, exons 2-89 and 11-10
4643	14531	24324	1.14	2.5E-01	Q27225 AF007768.1	SWISSPROT	MOLT-INHIBITING HORMONE PRECURSOR ((MIH) Choristoneura fumiferana diapause associated protein 2 (DAP2) mRNA, complete cds
4672 4698 4726	11 11	24351	3.16	2.5E-01 2.5E-01 2.5E-01		NT NT EST_HUMAN	Vibrio cholerae chromosome II, section 73 of 93 of the complete chromosome Mus musculus annexin V gene, intron 4 segment containing 5' LTR and gag portion of MuERV-L (murine endogenous retrovirus) element 601437468F1 NIH_MGC_72 Homo sapiens cDNA, done IMAGE:3922600 5'

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
4760	14635	24421	0.89	2.5E-01 A	B011070.1	TN	Mus musculus gene for uncoupling protein 3, 5'-itanking region and partial 5UTR
5169	15035	24802	0.86	2.5E-01	AW663183.1	EST_HUMAN	hh75f09.y1 NCI_CGAP_GU1 Homo sapiens cDNA clone IMAGE:2968649 5' similar to contains TAR1.t2 TAR1 repetitive element;
5169	15035	24803	0.86	2.5E-01	2.5E-01 AW663183.1	EST HUMAN	hh75f09.y1 NCI_CGAP_GU1 Homo sapiens cDivA clone IMAGE:2968649 5' similar to contains TAR1.t2 TAR1 repetitive element;
5264	15186		11.62	2.5E-01 S	83390.1	NT	T3 receptor-associating cofactor-1 [human, fetal [iver, mRNA, 2930 nt]
8366	16229		1.32	2.5E-01 A	F134119.1	LN LN	Mus musculus SKD1 (Skd1) gene, complete cds
6500	16359	26532	3.73	2.5E-01 A	163282.2	LN	Homo sapiens chromosome 21 segment HS21C082
6583	16463	26655	2.99	2.5E-01 Bi	BF109040.1	EST_HUMAN	7157a03.x1 Scares_NSF_F8_9W_OT_PA_P_S1 Homo sapiens cDNA clone IMAGE:3525389 3'
6788	16687	26858	2.25	2.5E-01	BF038595.1		601459238F1 NIH_MGC_66 Homo sapiens cDNA clone IMAGE:3862809 5'
7020	16897	27087	3.95	2.5E-01	H53236.1	T_HUMAN	yq84f07.r1 Soares fetal liver spleen 1NFLS Home saplens cDNA clone IMAGE:202501 5'
7481	17351	27554	16.11	2.5E-01	U89651.2	NT	Homo sapiens matrix metalloprotalnase MMP Raisl-1 gene, promoter region
7481	17351	27555	18.11	2.5E-01	89651.2	NT	Homo sapiens matrix metalloproteinase MMP Raisl-1 gene, promoter region
7522	17341	27547	2.04	2.5E-01 A	F085164.1	NT	Hordeum vulgare receptor-like kinase LRK10 gene, partial cds
7522	17341	27548	2.04	2.5E-01		NT	Hordeum vulgare receptor-like kinase LRK10 gene, partial cds
7814	17684	27904	1.5	2.5E-01	2.5E-01 AW581997.1	EST_HUMAN	RC3-ST0186-130100-015-a07 ST0186 Homo saplens cDNA
8010	17860	28105	1 82	2 5F-01	AW152246 1	FST HIMAN	xg40c10.x1 NCI_CGAP_Ut1 Homo sapiens cDNA clone IMAGE:2630034 3' similar to contains Alu repetitive element contains Alu repetitive element contains element MSR1 repetitive element :
2	17861	28106	1,68	2.5E-01	X58491.1	L	Mause L1Md LINE DNA
8426	18300	28558	2.32	2.5E-01	D50914.1	LN	Human mRNA for KIAA0124 gene, partial cds
9074	18851	29117	2.45	2.5E-01	AF200528.1	LN L	Zea mays cellulose synthase-4 (CesA-4) mRNA, complete cds
9100	19729		4.2	1	2.5E-01 AL161541.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 41
6228	19581	25072	1.28		AF170072.1	INT	Spodoptera frugiperda CALNUC mRNA, complete cds
541	10482	20282	1.53		AA936316.1	EST_HUMAN	on70d04.s1 Soares_NFL_T_GBC_S1 Homo sepiens cDNA clone IMAGE:1562023 3'
830	10757	20908	2.38		2.4E-01 BF576124.1	EST_HUMAN	602132442F1 NIH_MGC_81 Hamo saplens cDNA clone IMAGE:4271578 5'
1282	11190	21041	17.41	2.4E-01	2.4E-01 AJ289880.1	NT	Homo sapiens KIAA0851 gene (partial), XT3 gene and LZTFL1 gene
1282	11190	21042	17.41	2.4E-01	AJ289880.1	NT	Homo sapiens KIAA0851 gene (partial), XT3 gene and LZTFL1 gene
1362	11268	21123	1.04		2.4E-01 Y17293.1	NT	Homo sapiens FLI-1 gene, partial
1808	11705		24.08	2.4E-01	AF267753.1	NT	Mesembryanthemum crystallinum putativa potassium channel protein Mkt1p mRNA, complete cds
1858	11754	21629	1.41	2.4E-01	K I	NT	Zaccys dhumnades fructose-1,6-bisphosphatase inRNA, complete cds
2091	11980	21875	0.88	2.4E-01	AF111168.2	N	Homo sapiens serine palmitoyl transferase, subunit II gene, complete cds; and unknown genes
2122	' 1		1.16	24E-01 P	45384	SWISSPROT	IMMUNOGLOBULIN A1 PROTEASE PRECURSOR (IGA1 PROTEASE)
82	12101	22005	. 2.01	2.4E-01 A	E000680.1	LN	Aquifex aeclicus section 12 of 109 of the complete genome

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Table 4
Single Exon Probes Expressed in Heart

Top Hit Descriptor	7h23d04x1 NCI_CGAP_Co16 Homo sapiens cDNA clone IMAGE:3316807 3' similar to SW:PRSB_XENLA 042588 28S PROTEASE REGULATORY SUBL'NIT 6A ;	D.discoldeum (Ax3-K) ponA gene	S.pombe swi6 gene	Bovine adenovirus 3 complete genome	Oryza longistaminata receptor kinase-like protein, famlly member D, and retrofit (gag/pol) genes, complete cds	H.sapiens AGT gene, Pstl fragment of intron 4	QV1-HT0412-020400-136-b10 HT0412 Homo saplens cDNA	Hepatitis C virus genomic RNA for polyprotein, complete cds	Mus musculus Wrn protein (Wrn) gene, complete cds	Mus musculus Wrn protein (Wrn) gene, complete cds	7154404.x1 NCI_CGAP_Br16 Homo saplens cDNA clone IMAGE:3338503 3' similar to SW:SFR4_HUMAN Q08170 SPLICING FACTOR, ARGININE/SERINE-RICH 4 ;contains element TAR1 TAR1 repetitive element		Drosophila melanogaster p38a MAP kinase gene, complete cds	Homo sapiens HSPC142 protein (HSPC142), mRNA	wc62c11.x1 NCI_CGAP_Pan1 Homo sapiens cDNA clone IMAGE:2323220 3' similar to gb:J03464 PROCOLLAGEN ALPHA 2(1) CHAIN PRECURSOR (HUMAN);	Bos taurus guanylyi cyclase-activating protein 2 (guca2) mRNA, complete cds	Tetrahymena thermophila macronuclear gene encoding ribosomal protein L3, exons 1-2	wd43e02x1 Sogres_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2330906 3' similar to contains	COLLAGEN ALPHA 1(X) CHAIN PRECURSOR	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 8	Mus musculus type 1 sigma receptor gene, compliate cds	P.asiatioa mosaic virus genomio RNA	Homo sapiens fragile 16D oxido reductase (FOR) gene, exon 6	Arabidopsis thaliana ethylene-insensitive3-iike1 (EIL1) mRNA, complete cds	Mus musculus mRNA for putative mc7 protein (mc7 gene)	Gallus gallus gene coding for a-actin	RC3-CT0413-100800-023-b06 CT0413 Homo saplens cDNA	Homo saplens chromosome 21 segment HS21C081	aromatase [Poephila guttata≂zebra finches, ovary, mRNA, 3188 nt]
Top Hit Database Source	EST_HUMAN			I	뒫	F	EST_HUMAN	IN		I		EST_HUMAN	IN IN			N	Į.	144	SWISSPROT		F					Į,	EST_HUMAN	NT	
Top Hit Acession No.	BF002171.1	236534.1	X71783.1	AF030154.1	U72726.1	X74209.1	BE160080.1	D00944.1	AF091216.1	AF091216.1		BF592336.1	AF035546.1	7661801 NT	AI698989.1	L43001.1	AJ012585.1	A 100014	Albessers.1	AL161494.2	AF030189.1	221647.1	AF217491.1	AF004213.1	AJ278191.1	V01507.1	BF229975.1	AL163281.2	S75898.1
Most Similar (Top) Hit BLAST E Value	2.4E-01	2.4E-01	2.4E-01	2.4E-01	2.4E-01	2.4E-01	2.4E-01	2.4E-01	2,4E-01	2.4E-01		2.4E-01	2.4E-01	2.4E-01	2.4E-01	2.4E-01	2.4E-01	L	2.4E-01	2.4E-01	2.4E-01		2.4E-01	2.4E-01	2.4E-01	2.4E-01	2.4E-01	2.4E-01	2.3E-01
Expression Signal	0.85	1.63	2.14	9.68	2.82	1.85	0.88	50.15	7.53	7.53		2.13	2.66	2.15	1.79	8.87	1.62	, i	9.72	3.63	1.99	2.42	1.55	2.39	2.64	1.69	1.26	3.35	6.0
ORF SEQ ID NO:	22111	22260			•	22831	24486	24622	25451			25583	25648	25714	25968				2/511		28347		29109						20153
Exon SEQ ID NO:	12213	ļ		12618	13020	13035	14700	14856	15389	15389		15508	15555	15613	15842	l	1	i	17815	_	L	L	L	19526	18946	19509	19716	L	10330
Probe SEQ ID NO:	2332	2491	2734	2758	3083	3109	4817	4981	5469	5469		5593	5642	5705	5937	6345	0989	;	7985	8149	8209	8534	9030	9162	9222	9439	9650	9865	383

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Table 4
Single Exon Probes Expressed in Heart

1		\neg		r—	г	т-	_				Т	-			٦									Т	Т	Τ-	Т	72
	Tap Hit Descriptor	C.familiaris rom1 gane	as27e12.x1 Barstead aorta HPLRB6 Homo saplens cDNA clone IMAGE:2318446 3' similar to gb:X13238 CYTOCHROME C OXIDASE POLYPEPTIDE VIC PRECURSOR (HUMAN);	as27e12.X1 Barstead acrta HPLRB6 Homo sapiens cDNA clone IMAGE:2318446 3' similar to gb:X13238 CYTOCHROME C OXIDASE POLYPEPTIDE VIC PRECURSOR (HUMAN);	as42f12.x1 Barstead aorta HPLRB6 Homo sapiens cDNA clone IMAGE:2319887 3' similar to contains Alu repetitive element,	Glycine max resistance protein LM17 precursor RNA, partial cds	Mus musculus myosin XV (Myo15), mRNA	601511573F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3912859 5'	za12e08.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:292358 5'	Oxydricha nova macronuclear telomere-binding protein alpha subunit (tel-elpha alanine version) gene, complete cds	Haemophilus influenzae genes for HinclI restriction-modification system (HinclI methyltransferase (EC	2.1.1.72) and Hincll endonuclease (EC 3.1.21.4))	MR0-HT0559-240400-014-g11 HT0559 Homo sapiens cDNA	Rhizobium leguminosarum partial genomic DNA for exopolysaccharide biosynthesis genes	601646155R2 NIH_MGC_59 Homo sapiens cDNA clone IMAGE:4102092 3'	Mus musculus partial mRNA for muscle protein 634 (mg534 gene)	Mus musculus partial mRNA for muscle protein 634 (mg534 gene)	Chlamydophila pneumoniae AR39, section 4 of 94 of the complete genome	Borrella burgdorferi 2.9-6 locus, ORF-A-D genes complete cds and REP+ gene, partial cds	HCOEST44 HT29M6 Homo sapiens cDNA clone HCoE44 5'	chn1424.seq.F Human fetal heart, Lambda ZAP Express Homo saplens cDNA 5'	PM4-SN0012-030400-001-b06 SN0012 Homo stiplens cDNA	xx21d07.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2813773 3' similar to TR:Q9Z175 Q9Z175LY3YL OXIDASE-RELATED PROTEINI2: contains PTR5.b2 TAR1 repetitive element :	601507202F1 NIH_MGC_71 Home sapiens cDNA clone IMAGE:3908689 5'	602144459F1 NIH_MGC_48 Homo sapiens cDNA clone IMAGE:4297719 5'	Rattus norvegicus mRNA for acid gated ion channel	Rattus norvegicus mRNA for acid gated ion channel	nao39h12.x1 Lupski_sciatio_nerve Homo sapiens\oDNA clone \MAGE:3395950 3' similar to contains element MER38 repetitive element :
2.6	Top Hit Database Source	NT	EST_HUMAN	EST_HUMAN	EST HUMAN	NT	R	EST_HUMAN	EST_HUMAN	Į.		TN	EST_HUMAN	NT	EST_HUMAN	TN	IN	TN	TN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	N	N	EST HUMAN
	Top Hit Acession No.	X96587.1	AI708840.1	AI708840.1	AI718148.1	AF175389.1	6754779 NT	BE888071.1	N80983.1	M68931.1		X52124.1	BE173060.1	AJ283261.1	BF133577.1	AJ250189.1	AJ250189.1	AE002167.2	U45428.1	127231.1	AA089819.1	AW863940.1	AW303623.1	BE882464.1	BF663319.1	AJ006519.1	AJ006519.1	8F475611.1
	Most Similar (Top) Hit BLAST E Value	2.3E-01	2.3E-01	2.3E-01	2.3E-01	2.3E-01	2.3E-01	2.3E-01	2.3E-01	2.3E-01		2.3E-01	2.3E-01	2.3E-01	2.3E-01	2.3E-01	2.3E-01	2.3E-01	2.3E-01	2.3E-01	2.3E-01	2.3E-01	2.35-01	2.3E-01	2.3E-01	2.3E-01	2.3E-01	2.3E-01
	Expression Signal	4.83	1.87	1.87	3.93	2.62	3.19	1.59	2.9	2.28		1.37	2.54	2.26	4.95	284	2.84	2.39	2.53	20.48	1.65	2.07	2.05	4.88	1.93	2.09	4.54	2.67
	ORF SEQ ID NO:	25361		25618				26519		26741	L		27787	27811		28863	28664	28830					24990					
	Exon SEQ ID NO:	15314	15533	15533	16005	16246	16345	16349	16418	16544		17507	17562	17590	17824	18397	18397	18547	18898	18951	19477	18970	19665	19708	19060	19088	19088	1
	Probe SEQ ID NO:	5385	5618	5618	6111	6384	6487	6491	929	6664		7657	7712	7740	7974	8525	8525	8658	9144	9232	9528	9566	7686	9358	9407	9456	9549	9793
,		_					_	_	_		_	_	_		_	_	-	_	_	_	_			-		_		

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	Top Hit Descriptor	oz14a10.x1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:1675290 3' similar to TR:Q13040 Q13040 ATP-BINDING CASSETTE PROTEIN ;	Homo sapiens PPAR delta gene, promoter región	Trimeresurus malabaricus cytb gene, partial cds; mitochondrial gene for mitochondrial product	Fresh-water sponge Emf1 alpha collagen (COLi;1) gene	602085608F1 NIH_MGC_83 Homo saplens cDNA clone IMAGE:4249969 5'	601462629F1 NIH_MGC_67 Homo sapiens cDNA clone IMAGE:3866190 5'	601462829F1 NIH_MGC_67 Homo sapiens cDNA clone IMAGE:3866190 5'	PM2-HT0353-281299-003-a12 HT0353 Homo sapiens cDNA	PM2-HT0353-281299-003-e12 HT0353 Homo sapiens cDNA	Homo sapiens FRA3B common fragile region, diadenosine triphosphate hydrolase (FHIT) gene, exon 5	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 62	Xiphophorus maculatus truncated Rex1 retrotransposon reverse transcriptase (RT) pseudogene	Mus musculus ATP-binding cassette protein (Abcb8) mRNA, partial cds	Drosophila melanogaster UNC-119 (unc-119) gene, complete cds	Mus musculus mixed lineage kinase 3 (Mik3) and two pore domain K+ channel subunit (Konk6) genes,	complete cas	Mus musculus MAP kinase kinase 1 (Mekk1) mRNA, complete cds	Mus musculus MAP kinase kinase kinase 1 (Mekk1) mRNA, complete cds	Human scRNA (BC200 beta) pseudogene	Human scRNA (BC200 beta) pseudogene	B.abortus bp26 gene	Human beta-cytoplasmic actin (ACTBP9) pseudogene	zq87c05.r1 Stratagene hNT neuron (#937233) Homo sapiens cDNA clone IMAGE:648968 5'	Human dystrophin gene	Mus musculus vinculin gene, exon 3	Homo sapiens diaphanous (Drosophila, homolog) 2 (DIAPH2), transcript variant 156, mRNA	Synechocystis sp. PCC6803 complete genome, 19/27, 2392729-2538999	AV756238 BM Homo sapiens cDNA clone BMFAHC06 5'	Human glycophorin B gene, exon 4	Human glycophorin B gene, exon 4	Mus musculus nm23-M1 gane, promoter region	Thermotoga maritima section 25 of 136 of the complete genome
וופוס בעמון ו ומסק מופויו	Top Hit Database Source	EST_HUMAN TR	NT		NT		EST_HUMAN 60			EST_HUMAN PA	<u> </u>	NT	文	NT	Į.						H LN		Ή LN	EST_HUMAN Z	H L	INT IN			T_HUMAN	NT	NT		TN
	Top Hit Acesslon No.	A1052190.1	AF187850.1	AF171901.1	M34640.1	2.2E-01 BF677538.1	BE618258.1	BE618258.1	BE155625.1	BE155625.1	AF020503.1	AL161562.2	AF155728.1	AF213391.1	AF119102.1		۹.	ч.	AF117340.1	J01307.1	J01307.1	254148.1	50604.1	AA211216.1	M86524.1	L13289.1		D84000.1	AV756238.1	M24136.1		AF155143.1	AE001713.1
	Most Similar (Top) Hit BLAST E Value	2.2E-01	2.2E-01	2.2E-01	2.2至-01	2.2E-01	2.2E-01 BE	2.2至-01	2.2€-01	2.2€-01	2.2E-01	2.2年-01	2.2E-01	2.2E-01	2.25-01		2.2E-01	2.2E-01	2.2E-01	2.2至-01	2.2E-01	2.2E-01	2.2E-01	2.2至-01	2.2E-01	2.2E-01	2.2€-01	2.2E-01	2.2E-01	2.2€-01	2.2E-01	2.2E-01	2.2E-01
	Expression Signal	0.98	3.13	0.91	2.78	6.24	2.41	2.41	4.04	4.04	1.59	2.67	1.05	0.81	1.19		5.07	1.97	1.97	1.18	1.16	1.09	1.22	2.47	1.19	1.2	1.71	3.99	10.59	2.01	2.01	2.19	4.27
	ORF SEQ ID NO:	19885	21311		21829	22131	22307	22308	22570	22571				23688				23840			23934			24396	24549		25495				26343		72272
	Exon SEQ ID NO:	10068	11450	11865	11833		12417	12417	12781	12781	12817	<u>l</u> _	13655	13913	14018	ļ		14066			14157	14294	14605	14610	14771	14850	L	I	16074	16182	16182	16556	17034
	Probe SEQ ID NO:	8	1545	1972	2042	2354	2543	2543	2853	2853	2890	3346	3743	4007	4118		4125	4166	4166	4258	4258	4399	4719	4724	4891	4975	5513	5518	6189	6319	6318	6676	7157

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											Γ		.A12	actin														Γ				
Top Hit Descriptor	PM3-CT0263-241299-009-b07 CT0263 Homo sapiens cDNA	Mus musculus deformed epidermal autoregulatory factor 1 (Drosophila) (Deaff), mRNA	WR1-TN0045-110900-006-c02 TN0045 Homo sapiens cDNA	za04f08.r1 Soares melanocyte 2NbHM Homo sapiens cDNA clone IMAGE:291591 5'	LARGE PROLINE-RICH PROTEIN BAT2 (HLA-B-ASSOCIATED TRANSCRIPT 2)	Brachydanio rerio ependymin beta and gamma chains (Epd) gene, complete cds	Funaria hygrometrica chloroplast-localized small heat shock protein (CPsHSP21) mRNA, complete cds;	nuclear gene for chloroplast product	601869724F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:4100189 5	Drosophila 68C glue gene cluster	Homo saplens H-2K binding factor-2 (LOC51580), mRNA	601446957F1 NIH_MGC_65 Homo sapiens cDNA clone IMAGE:3850670 5'	Homo sepiens chromosome Xq28 melanoma antigen family A2a (MAGEA2A), melanoma antigen family A12	(MAGEA12), melanoma antigen family A2b (MAGEA2B), melanoma antigen family A3 (MAGEA3), caltractin	(CALT), NAD(P)H dehydrogenase-like protein (NSDHL), and LI>	Vitis vinifera cultivar Pinot Noir plasma membrano aquaporin (PIP1a) mRNA, complete cds	RC1-CT0249-141199-021-g04 CT0249 Homo sapiens cDNA	AV694801 GKC Homo sapiens cDNA clone GKCAHB02 5'	nm31e11.s1 NCI_CGAP_Lip2 Homo sapiens cDNA clone IMAGE:1061804	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 16	Chlamydia muridarum, section 45 of 85 of the complete genome	Mus musculus interferon (alpha and beta) receptor 2 (Ifnar2), mRNA	Mus musculus interferon (alpha and beta) receptor 2 (ifmar2), mRNA	ok73e02.s1 NCI_CGAP_GC4 Homo sepiens cDNA done IMAGE:1519610 3' similar to gb:K02765 COMPLEMENT C3 PRECURSOR (HUMAN);	602083129F1 NIH_MGC_81 Homo sapiens cDNA clone IMAGE:4247503 5	Homo sepiens potassium voltage-gated channel, subfamily H (eag-related), member 4 (KCNH4), mRNA	Beta vulgaris mitochondrion, complete genome	IMMEDIATE EARLY PROTEIN IE180	IMMEDIATE-EARLY PROTEIN IE180	Homo saplens mRNA for KIAA1215 protein, partiel cds	Homo sapiens pshsp47 gene, complete cds	Homo sapiens hox11 proto-oncogene, exons 1 to 5 and hug-1 gene
Top Hit Database Source	EST_HUMAN	N.	EST_HUMAN	EST_HUMAN	SWISSPROT	ΝΤ		LZ.	EST_HUMAN	LN	N	EST_HUMAN			L	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	TN	N _T	ΙN	NT L	EST HUMAN	EST_HUMAN	Ė	Z	SWISSPROT	SWISSPROT	NT	TN	NT
Top Hit Acession No.	AW855039.1	8393247 NT	BF376354.1	W02988.1	P48634	M89643.1		AF197941.1	BF206507.1	X01918.1	7706216 NT	BE870959.1			J82671.2	AF188843.1	4W361098.1	AV694801.1	AA569289.1	AL161504.2	AE002314,2	6754299 NT	6754299 NT	AA906824.1	BF695073.1	6912445 NT	D838361 NT	P11675	P11675	AB033041.1		AJ009794.1
Most Similar (Top) Hit BLAST E Value		2.2E-01			22至-01							2.2€-01		-			2.2E-01	2.2€-01		2.1E-01	2.15-01	2.1E-01	2.1E-01		2.1E-01	2.15-01	2.1E-01	2.1E-01	_			2.1E-01
Expression Signal	2.29	1.68	1.39	4.1	13.13	3.98		3.67	22	4.94	2.91	2.2			3.72	2.34	2.58	3.75	1.74	6.0	2.18	1.24	1.24	1.84	3.08	1.65	5.05	1.01	1.01	1.28	1.21	1.26
ORF SEQ ID NO:		27341	27382	27421	27550			27720	27794	28824	28149						24887		20727	20729		20936	20937	21642	21899	22813		23651	23662			24439
Exon SEQ ID NO:	17094	17147				17315		- 1	17569	18540	17905	18854					15093	19713	10879	10881	11023	11091	11091	11767	12000	12821	L	L	L.		i	14651
Probe SEQ (D NO:	7217	7270	7304	7353	7366	7397		7648	7719	8723	8766	8077			9183	9269	9379	9875	985	828	1108	1180	1180	1871	2111	2894	3736	3967	3967	4279	4474	4766

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	Top Hit Descriptor	Saccharomyces cerevisiae tau 138 (TFC3) gene, complete cds	602152001F1 NIH_MGC_81 Homo sapiens cDNA clone IMAGE:4283001 5	Human offactory receptor (OR17-2) gene, partial cds	Archaeoglobus fulgidus section 135 of 172 of the complete genome	Canis familiaris keratin (KRT9) gene, complete cds	Glycine max malate dehydrogenase (Mdh-2) gene, nuclear gene encoding mitochondrial protein, partial cds	Glycine max malate dehydrogenase (Mdh-2) gene, nuclear gene encoding mitochondrial protein, partial cds	Mus musculus erythrocyte protein band 4.1-like 3 (Epb4.1B), mRNA	Haemophilus influenzae hmcD, putative haemocin processing protein (hmcC), putative ABC transporter (hmcB), putative haemocin structural protein (hmcA), and haemocin immunity protein (hmcI) genes, complete	cds	S.cerevisiae chromosome II reading frame ORF YBL025w	A,thaliana mRNA for AtRanBP1b protein	Homo saplens p53R2 gene for ribonucleotide reductase, exon 6	Beta vulgaris mRNA for elongation factor 1-beta	DIACYLGLYCEROL KINASE, DELTA (DIGLYCERIDE KINASE) (DGK-DELTA) (DAG KINASE DELTA) (80 KD DIACYLGLYCEROL KINASE)	Homo sapiens pancreatic polypeptide 2 (PPY2), mRNA	RC3-HT0622-040500-013-b11 HT0622 Homo sapiens cDNA	Homo saplens fragile 16D oxido reductase (FOR) tiene, exons 8, 9, and partial cds	Нитап дтапијіп деле	7a59e02x1 NCI_CGAP_GC6 Homo sapiens cDNA clone IMAGE:3223034 3'	Salvelinus alpinus mitochondrion, complete genomie	Gallus gallus mRNA for avena, complete cds	Homo saplens CGI-18 protein (LOC51008), mRNA	O.cunniculus germline igH heavy chain V-H pseudogene, allotype VHa2	Mus musculus Major Histocompatibility Locus class II region	Synechocystis sp. PCC8803 complete genome, 7/27, 781449-920915	Homo saplens chromosome 21 segment HS210013	Homo saplens rac1 gene	PM1-HT0422-291299-002-c06 HT0422 Homo saplens cDNA	Homo sapiens dystrobrevin, alpha (DTNA), mRNA
10 - 10 - 10 - 10 - 10 - 10 - 10 - 10 -	Top Hit Detabase Source	NT.	EST_HUMAN	NT	NT	NT	NT	TN	ᅜ		NT	NT	NT	IN	LN	SWISSPROT	NT L	EST_HUMAN	TN	IN	EST_HUMAN	TN	IN	TN	IN	IN	LN	NT	NT	EST_HUMAN	NT
	Top Hit Acession No.	M98261.1	BF672695.1	J U04642.1	AE000972.1	AF000949.1	AF068687.1	AF068687.1	7305030 NT		U68399.1	Z35786.1	X97378.1	9.1	Z97067.1	P52824	11036647 NT	BE180422.1	AF217490.1		BE672330.1	5835904 NT	AB017437.1	7705601 NT	M77085.1	AF027865.1	D90905.1	1 AL163213.2	2.0E-01 AJ132695.5	2.0E-01 AW384937.1	4503408 NT
	Most Similar (Top) Hit BLAST E Value	2.1E-01	2.1E-01	2.1E-01	2.1E-01	2.1E-01	2.1E-01		2.1E-01	٠			2.1E-01	2.1E-01	2.1E-01	2.1E-01 P52824	2.1E-01		2.1E-01	2.1E-01	2.1E-01	2.1E-01	2.0E-01	2.0E-01	2.0E-01	2.0E-01	2.05-01	2.0E-01	2.0E-01	2.0E-01	2.0E-01
	Expression Signal	66.0	5.89	1.86	1.97	1.74	1,35	1.35	1.21		4.78	6.88	2.38	1.19	2.49	1.49	2.31	2.15	1.6	1.4	1.29	1.29	1.72	2.39	1.19	1.81	1.03	2.57	1.42	1.29	13.51
	ORF SEQ ID NO:		24938			26572	26598	26597				27289	27553					28964			25174		19983			20563				21047	
	Exon SEQ ID NO:		i			16393	16417	16417	16585		16829	17101	17349		17767	ļ	ļ	18674	Į	19634	19382	19436	10165		10817	10722	10915			1	11376
	Probe SEQ ID NO:	5134	5243	6123	6385	6535	6559	6559	6205		6951	7224	7479	7547	7917	7929	8840	8862	9522	9730	9905	9979	<u>8</u>	523	684	783	982	1109	1234	1286	1471

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
1534	11438		2.51	2.0E-01	2.0E-01 AB007974.1	NT	Homo sapiens mRNA, chromosome 1 specific trenscript KIAA0505
1538	11442	21300		2.0E-01	2.0E-01 AF260700.1	NT	Homo sapiens sodium/todide symporter mRNA, partial cds
1688	11590		1.82	2.0E-01	2.0E-01 AF111170.3	TN	Homo saplens 14q32 Jagged2 gene, complete cds; and unknown gene
1723	11624		1.64		U67525.1	NT	Methanococcus Jannaschii section 67 of 150 of the complete genome
1847	11743	21618				EST_HUMAN	601449441F1 NIH_MGC_65 Homo septens cDNA done IMAGE:3853330 5
1847	11743	21619	1.33			EST_HUMAN	601449441F1 NIH_MGC_65 Homo sepiens cDNA clone IMAGE:3853330 5
523	12181		1.67		2.0E-01 X82877.1	님	H.sapiens Na≁-D-glucose cotransport regulator gene
2861	12789		9:00	2.0E-01		IN	Homo saptens full tength insert cDNA YH85A11
3442	13359	23168	80		P46607	SWISSPROT	HOMEOBOX PROTEIN GLABRA2 (HOMEOBOX-LEUCINE ZIPPER PROTEIN ATHB-10) (HD-ZIP PROTEIN ATHB-10)
							xp15b02.x1 NCI_CGAP_HN9 Homo saplens cDNA clone IMAGE:2740395 3' similar to contains element
3520	13436		0.89	2.0E-01	2.0E-01 AW238005.1	EST_HUMAN	MER21 repetitive element;
3646	13560	23348	0.81	2.0E-01 P34841	P34641	SWISSPROT	CED-11 PROTEIN
3945	13853	23628	9.0	2.0E-01	2.0E-01 X83997.1	F	C, parasitica eapC gene
4484	14358		8.47	2.0E-01	2.0E-01 BE826165.1	EST_HUMAN	QV4-EN0032-190500-223-e03 EN0032 Homo sablens cDNA
4859	14739	24519	1.07	2.0E-01	2.0E-01 AF147083.1	NT	Homo sapiens gamma-glutamy/ hydrolase gene, exons 8 and 9 and complete cds
4972	14847	24618	6.07	2.0E-01	8922080 NT	NT	Homo sapiens hypothetical protein ASH1 (ASH1), mRNA
5041	14913	24687	1.1	2.0E-01 Y	Y19216.1	NT	Homo sapiens putative psihHbD pseudogene for hair keralin, exons 1 to 9
5342	15263	25089	2.55	2.0E-01	96600	NT	Rat SOD-2 gene for manganese-containing superoxide dismutase
5510	15428				11432540 NT	INT	Homo saplens dual oxidase-like domains 2 (DUOX2), mRNA
5694	15603	25705	5.29		U15300.1	NT	Saccharomyces cerevisiae Hal5p (HAL5) mRNA, complete cds
5874	15780				31033.1	NT	M.auratus mu class glutathione transferase gene
5934	15839	25962		2.0E-01		EST_HUMAN	PM1-CT0247-141099-001-g08 CT0247 Homo sapiens cDNA
6837	16517		6.95	2.0E-01	F028026.1	NT	Andes virus strain Ol23133 glycoprotein G1 and G2 precursor, gene, partial cds
6772	16851	26839				NT	M.musculus scp2 gene exon 14
7511	17299		4.39			NT	Chlamydia trachomatis section 5 of 87 of the complete genome
7692	17542		2.07		2.0E-01 AF146692.1	NT	Homo sapiens filamin 2 (FLN2) mRNA, complete cds
7774	17624				2.0E-01 AF086907.1	NT	Arabidopsis thaliana root gravitropism control protein (PIN2) gene, complete cds
7774	17624	27858			2.0E-01 AF086907.1	NT	Arabidopsis thaliana root gravitropism control protein (PIN2) gene, complete cds
8214	18098	28350	2.7	2.0E-01		NT	Salvelinus pluvius mRNA for transferrin, complete ods
8214	18098			2.0E-01		NT	Salvelinus pluvius mRNA for transferrin, complete cds
8503	19118	i			2.0E-01 AF208637.2	뒫	Pimephales promelas liver glucose-6-phosphate-1-dehydrogenase mRNA, partial cds
9747	19308	25202			2.0E-01 AI023592.1	EST_HUMAN	ov80a10.s1 Soares_testis_NHT Homo saplens cDNA clone IMAGE:1643810 3'
9769	19288		6.61	2.0E-01	2.0E-01 AF078164.2	닐	Homo sapiens Ku70-binding protein (KUB3) mRNA, partial cds

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Table 4
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Probe SEQ ID NO:	Exen SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acesslon No.	Top Hit Database Source	Top Hit Descriptor
9897	19374	25193	1.6	2.0E-01	11528495 NT	TN	Mus musculus fructosamine 3 kinase (Fn3k), mFNA
108	10088		10.35	1.9E-01	7549743 NT	NT	Rattus norvegicus Aryl hydrocarbon receptor nuclear translocator 1 (Arnt1), mRNA
349	10308	20126	9	1.9E-01	AF004353.1	NT	Mus musculus pale ear (ep) gene, wild type allele, 3' region, partial cds
640	10577	20392	1.31	1.9E-01	U32581.2	TN	Homo sapiens lambda/iota protein kinase C-interacting protein mRNA, complete cds
640		20393	1.31	1.9E-01	U32581.2	NT	Homo saplens lambda/iota protein kinase C-interciting protein mRNA, complete cds
647	10584	20400	5.37	1.9E-01	BE070801.1	EST_HUMAN	RC3-BT0502-251199-011-d01 BT0502 Homo sapiens cDNA
648	10584	20400	7.32	1.9E-01	BE070801.1	EST_HUMAN	RC3-BT0502-251199-011-d01 BT0502 Homo saplens cDNA
970	10893		1.61	1.9E-01	T305180 NT		Mus musculus interleukin 2 receptor, gemma chain (II2rg), mRNA
1088	11004	20846	9.3	1.9E-01	AA358813.1	EST_HUMAN	EST67784 Fetal lung II Homo saplens cDNA 5' end
1348		21110	2.3	1.9E-01	AF081282.1	TN	Sorghum bicolor 22 kDa kafirin cluster
1416	11322		3.91	1.9E-01	AF184623.1	NT	Plasmodium vivax reticulocyte binding protein-2 (rbp-2) gene, complete cds
. 2330	12211	22109	3.31	1.9E-01	IN 892253	TN	Homo saplens hypothetical protein FLJ10581 (FLJ10581), mRNA
2892		22811	3.91	1.9E-01		TN	Sigmodon hispidus p53 gene, partial cds
2908	12834		5.55	1.9E-01	J00922.1	ΤΝ	Gallus gallus ovalbumin (Y) gene, complete cds
3349	13269	23072	3.4	1.9E-01	D13197.1	۲	Mouse gene for Immunoglobulin diversity region D1
3434	13351	23158	4.63		R16487.1	EST_HUMAN	yf42f10.r1 Scares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:129547 5'
3735	13647	23432	0.93	1.9E-01	AF264017.1	TN	Rattus norvegicus arylacetamide deacetylase genė, complete cds
3762	13675	23457	96'0		P39768	SWISSPROT	PAIR-RULE PROTEIN ODD-PAIRED
3910	13820	23600	3.02	1.9E-01	AB006784.1	F	Schizosaccharomyces pombe DNA for cytoplasmic dynein heavy chain, complete cds
3892	13899	23676	1.89	1.9E-01	AW754106.1	EST_HUMAN	CM3-CT0315-271199-045-b11 CT0315 Homo sapiens cDNA
4138	14038	23813	1.08	1.9E-01	BE834943.1	EST_HUMAN	MR1-FN0010-290700-007-d04 FN0010 Homo saplens cDNA
4369	14285	24049	68.0	1.9E-01	AL161493.2	LN.	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 5
4662	14548	24338	0.84	1.9E-01	293780.1	NT	Fugu rubripes genes encoding carbamoyi phosphate synthetase III, myosin light chain, MAP2
4912	14791	24566	0.86	1.9E-01	AW849203.1	EST_HUMAN	IL3-CT0215-180200-087-D02 CT0215 Homo saplens cDNA
4943	14821		1.04	1.9E-01	AF223642.1	TN	Rattus narvegicus chemokine receptor CXCR3 mRNA, complete cds
4962	14837	24605	1.11		095239	SWISSPROT	KINESIN-LIKE PROTEIN KIF4
5031	14903	24875	1.03	1.9E-01	AJ251178.1	FX	Phoca vitulina partial aar2B gene for alpha adrenergic receptor 2B
5113	14981	24755	0.99	1.9E-01	270296.1	ΝŢ	S.mansoni elastase HP1 gene
64.00	7,007		Ç	10 E 01	Ale31100 1	EST HIMAN	1593g12.X1 NCI_CGAP_GC6 Home sapiens cDNA clone IMAGE:2238886 3' similar to gb:M21574 ALPHA IDI ATEI ET.DERIVED GROWTH FACTOR RECEPTOR DRECHIRSOR (HUMAN)
5153	L	24789	66.0	L	79095	LN	Mus musculus Notch gene homolog 3, (Drosophila) (Notch3), mRNA
1444			86 7	l	AW/420440 4	EST HIMAN	7/29a07.x1 NCI_CGAP_Ut1 Homo sapiens cDNA clone IMAGE:2619444 3' similar to gb:M73779 RETINOIC ACID RECEPTOR AI PHA.1 (HIMAN):
	. 1		27.	3 10		NUMBER OF STREET	Harmon DNA releases one flat acted to the restole (DOI E1) ages 40
2466	200	25446	/0./	1.85-01	AF12/93/.1	2	India sapiers DIVA puyricaase epsilor catague submitty Ducir (COLET) Bere, exur ra

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Top Hit Descriptor	AU133116 NT2RP4 Homo sapiens cDNA clone NT2RP4001328 5'	yg09a12.s1 Soares infant brain 1NIB Homo sapiens cDNA clone IMAGE:31663 3' similar to contains MER13 repetitive element;	Arabidopsis thaliana serine/threonine protein phosphalase type one (TOPP8) gene, complete cds	Zea mays starch branching enzyme I (sbe1) gene, complete cds	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 57	Homo sapiens mRNA for KIAA1198 protein, partial cds	Arabidopsis thallana DNA chromosome 4, contig fragment No. 15	Arabidopsis thaliana DNA chromosome 4, contigi fragment No. 15	Homo sapiens calcium channel alpha/E subuni (CACNA/E) gene, exons 7-49, and partial cds, alternatively	opinosis. Homo sanians partial 5.HT4 recentor dena exorts 2 to 5.	Description programmes and the state of the	Orosophila magnogaster craftin light chair in his viv. Complete cus	Arabidopsis thaliana receptor-like kinase LECRK1 (LECRK1) gene, complete cds	Mus musculus p116Rip mRNA, complete cds	Mus musculus Cotg gene for chaperonin containing TCP-1 gamma subunit, partial cds	Homo sapiens calcium channel, voltage-dependent, beta 2 subunit (CACNB2) mRNA, and translated	products	Oryzias latipes gene for membrane guanylyl cyclase OIGC1, complete cds	wd71f02.x1 NCI_CGAP_Lu24 Homo saplens cDNA done IMAGE:2337051 3'	Dictyostellum discoideum plasmid Ddp5, complete genome	Yersinia pestis plasmid pCD1	Mus musculus guanylate nucleotide binding protein 1 (Gbp1), mRNA	Mus musculus guanylate nucleotide binding protein 1 (Gbp1), mRNA	Homo saplens latent transforming growth factor beta binding protein 4 (LTBP4) mRNA	qg22d10.x5 NCI_CGAP_Kid3 Homo sapiens cDNA ckne IMAGE:1761811 3' similer to TR:075936 075936 GAMMA BUTYROBETAINE HYDROXYLASE :	Mus musculus Scya6, Scya16-ps, Scya5 genes for small Inducible cytokine A6 precursor, small Inducible cytokine A9 precursor. Scya16 pseudodene, small inducible cytokine A5 precursor, complete cds	OVS. DT0018_081900_086_014 DT0018 Home cablene cDNA	Innovatifilm acarda EAFV broken (IEAFV) parts partial cde	24/a03.x1 Soares NFL T GBC S1 Homo eaplens cDNA clone IMAGE:2659756 31	QV0-BN0041-070300-147-c04 BN0041 Homo sapiens cDNA	
Top Hit Database Source	EST_HUMAN	EST HUMAN	TN	L	L N	LN N	LN	LN FN	<u> </u>					NT	NT		LN	LΝ	T_HUMAN	LΝ	LN		LN.		EST_HUMAN	Ŀ	T LI IMANI	NONCE	T HUMAN	Г	
Top Hit Acession No.	AU133116.1	R43212.1	U80922.1	AF072724.1	AL161557.2	AB033024.1	AL161503.2	AL161503.2	A E000001.4		Ţ		<u></u>	U73200.1	AB022090.1		2532	AB021490.2	A1912212.1	AF000580.1	AL117189.1	6753947 NT	6753947 NT	4505036 NT	AI733708.1	AB081807.1	Τ.]
Most Similar (Top) Hit BLAST E Value	1.9E-01	1.9E-01	1.9E-01		1.9E-01	1.9E-01	1.9E-01	1.9E-01	100						1.8E-01						1.8E-01	1.8E-01	1.8E-01	1.8E-01		т С	_			1.8E-01/	
Expression Signal	2.28	1.7	1.43	3.06	1.62	12.89	2.16	2.16	35	284		79.	1.28	2.28	1.22		2.41	2.15	0.85	1.21	5.28	1.28	1.29	0.92	1.83	, a	2 8	4.03	100	18	
ORF SEQ ID NO:		24863	26446	26478	26735	27109	28176	28177	26074	20080				19811	20041	1	20145	20493	20735	20832	21024	21252	21253			21644		+	22595	22803	
Exon SEQ ID NO:	15503	15119	16284	16310	16538	16918	17930	17930	2000	18788		19140	19548	10018	12663		10322	10661	10889	10980	11174	11392	11392	11700	11720	11780	42537	12704	12800	13012	
Probe SEQ ID NO:	5588	6162	6423	6448	6658	7041	8038	8038	0.07	0137	3 2	8240	9880	28	280		366	729	996	1075	1287	1487	1487	1803	1823	4874	2890	3000	2873	3083	

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					A		
Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
3570	13484	23275	1.07	1.8E-01	H03369.1	EST_HUMAN	1/45e01.s1 Soares placenta Nb2HP Homo sapiens cDNA clone IMAGE:1517043' similar to contains Alu repetitive element;
3570	13484	23276	1.07	1.8E-01	H03369.1	EST HUMAN	M45e01.s1 Soares placenta Nb2HP Homo sapiens cDNA clone IMAGE:151704 3' similar to contains Alu repetitive element,
4154	14054			1.8E-01	AJ271735.1	N.	Homo sapiens Xq pseudoautosomal region; segment 1/2
4238	14137		1.13	1.8E-01	D37954.1	NT	Bowne NB25 mRNA for MHC class II (BoLA-DQB), complete cds
4458	14350	24141	5.12	1.8E-01	AL161556.2	LN	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 58
4663	14549	24330	2.38	1 BF-01	AB051897 1	L Z	Mus musculus Scya6, Scya9, Scya16-ps, Scyaë genes for small Inducible cytokine A6 precursor, small Inducible cytokine A9 precursor. Scya16 pseudocene, small inducible cytokine A5 precursor, complete cds
4700	14586		0.83	1.8E-01	\sim	Ę	S.tuberosum mRNA for alcohol dehydrogenase
4977	14852	24618	1.77	1.8E-01	AW814270.1	EST_HUMAN	MR3-ST0203-151299-112-g06 ST0203 Homo saplens cDNA
5027	14800	24670	4.17	1.8E-01	AF181258.1	F	Mesocricetus auratus Na-taurochdate cotransporting polypeptide mRNA, partial cds
5053	14925	24697	1.28	1.8E-01	AI439881.1	EST_HUMAN	t67e04x1 NCI_CGAP_Lym12 Homo sapiens cDNA clone IMAGE:2134690 3'
5117	14985		1.03	1.8E-01	AJ000742.1	ΙN	Homo Sapiens hisH1 gene, 5' UTR
5549	15465	25535	1.41	1.8E-01	AL161594.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 90
5922	15827	25952	1.29	1.8E-01	Q9QY14	SWISSPROT	FORKHEAD BOX PROTEIN E3
5948	15853		2.5	1.8E-01	N94853.1	EST_HUMAN	yy62h02.r1 Soares_multiple_sclerosis_2NbHMSP Homo saplens cDNA clone IMAGE:278163 5
6179	16065	26214	1.38	1.8E-01	AB018561.1	NT	Citrullus lanatus mRNA for wsus, complete cds
6179	16085	28215	1.38	1.8E-01	AB018561.1	NT	Citrulius lanatus mRNA for wsus, complete cds
7382	17251	27458	1.72	1.8E-01	M73258.1	NT	Human cellular DNA/Human papillomavirus proviral DNA
7398			1.22	1.8E-01		INT	Bacteriophage Ike, complete genome
7933	17783	28022	1.19	1.8E-01	X63440.1	NT	M.musculus mRNA for P19-protein tyrosine phosphatase
8033	17925	28172	3.19	1.8E-01	X77336.1	NT	A.thaliana mRNA for ribonucleotide reductase R2
9908	17958	28205	99'9	1.8E-01	U38906.1	N.	Bacteriophage r1t integrase, repressor protein (rrc), dUTPase, holin and lysin genes, complete.cds
8118	16065	26214	2.9	1.8E-01	AB018561.1	N	Citrulius lanatus mRNA for wsus, complete cds
8118	16065	26215	2.9	1.8E-01		F	Citrulius lanatus mRNA for wsus, complete cds
8119			3.88	1.8E-01	AF019107.1	NT	Dictyostelium discoldeum unknown (DG1041) gene, complete cds
8381	18258	28507	2.64	1.8E-01	M59257.1	TN	Human carcinoembryonic antigen (OEA) gene, exon 4
8228	17907			1.8E-01	X57033.1	NT	B.taurus mRNA for potassium channel
8976	18781		2:83	1.8E-01	8394421		Rattus norvegicus Thromboxane receptor (Tbxa2r), mRNA
9169		25344		1.8E-01	BF348623.1	EST_HUMAN .	602019928F1 NCI_CGAP_Brn67 Hamo saplens ¢DNA clone IMAGE:4155318 5'
9825	19200		2.05	1.8E-01	Q96682	SWISSPROT	DNA TERMINAL PROTEIN (BELLETT PROTEIN) (PTP PROTEIN)

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression . Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
9738	19272		7.94	1.8E-01	1.8E-01 R24494.1	EST_HUMAN	yh48h10.r1 Soares placenta Nb2HP Homo sapions cDNA clone IMAGE:133027 5'
9779			1.63	1.8E-01	Y11114.1	NT	E.dispar mRNA for hexokinase (hxk1)
583	10503		1.8	1.7E-01	1.7E-01 BE385164.1	EST_HUMAN	801274604F1 NIH_MGC_20 Homo sapiens cDNA clone IMAGE:3615768 5
788	10717	20559	2.04	1.7E-01	1.7E-01 X53330.1	IN	P. dumerilli histone gene cluster for core histones H2A, H2B, H3 and H4
945	10870		1.89	1.7E-01	1.7E-01 P35616	SWISSPROT	NEUROFILAMENT TRIPLET L PROTEIN (NEWROFILAMENT LIGHT POLYPEPTIDE) (NF.L.)
1042	10960		1.6		1.7E-01 AF081810.1	NT	Lymantria dispar nucleopolyhedrovirus, complete genome
1042	10960	50803	1.6	1.7E-01	1.7E-01 AF081810.1	TN.	Lymantria dispar nucleopolyhedrovirus, complete genome
1938	11833		3.8	1.7E-01	1.7E-01 AF255051.1	NT	Homo sapiens BNIP3H (BNIP3H) gene, complete cds; nuclear gene for mitochondrial product
2829	12758	22548	1.93	1.7E-01	1.7E-01 AF000716.1	ĮN.	Vibrio cholerae hypoxanthine phosphoribosyltransferase (hpt) gene, partlal cds, hemagglutinin/protease regulatory protein (hapR) gene, complete cds, and YRAL VIBCO gene, partial cds
2829	12758	22549	1.83	1.7E-01	1.7E-01 AF000716.1	Ę	Vibrio cholerae hypoxanthine phosphoribosytransferase (hpt) gene, partial ods, hemagglutinin/protease regulatory protein (hapR) gene, complete ods, and YRAL VIBCO gene, partial ods
2896	1		1.74	1.7E-01	1.7E-01 AA336909.1	EST_HUMAN	EST41651 Endometrial fumor Homo sapiens cDNA 5' end
2967	12894		1.35	1.7E-01	1.7E-01 AJ238736.1	٦×	Naja naja atra ctx-1 gene, exons 1-3
2967	12894			1.7E-01	1.7E-01 AJ238736.1	TN	Naja naja atra ctv-1 gene, exons 1-3
3067	12994	22785	1.89	1.7E-01	1.7E-01 AF081514.1	NT	Taxus canadensis geranyigeranyi diphosphate synthase mRNA, complete cds
3401	13318	23119	1.96	1.7E-01	1.7E-01 AJ269505.1	TN	Anabasna sp. ORF4 (partial), ORF3, ORF2, ORF1, adpA gene, adpB gene, adpC gene, adpD gene, adpE gene and adpF gene
3557	13471	23262	1	1.7E-01	1.7E-01 AJ224877.1	IN	Homo sapiens hap1 gene, complete CDS
3859	13770	23562	4.41	1.7E-01	1.7E-01 AJ235377.1	ŊŢ	Homo sapiens derivative 11 breakpoint fragment: partial intron 10 of the ALL-1/MLL/HRX gene fused to intron 5 of the AF-4/FEL gene
4459	14353		1.88	1.7E-01	1.7E-01 X52936.1	TM	Schistocerca gregaria alpha repetitive DNA
4732	14817	24403	71	1 7E-01	1 7E-01 AI247635.1	EST HUMAN	qh57e09.x1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:1848808 3' sImilar to contains OFR.b1 OFR repetitive element :
4986	f .	L		1.7E-01	1.7E-01 U28376.1	N	Zea mays oalcium-dependent protein kinase (MZECDPK2) mRNA, complete cds
5067	1		1.18	1.7E-01	1.7E-01 AF072725.1	N	Zea mays starch branching enzyme IIb (ae) gene, complete cds
5321	15241	25046	1.76	1.7E-01	1.7E-01 AA470686.1	EST_HUMAN	ne13e02.s1 NCI_CGAP_Co3 Homo sapiens cDNA clone IMAGE:881066 3' similar to gb:M17886 60S ACIDIC RIBOSOMAL PROTEIN P1 (HUMAN);
5321	15241	25047	1.76		1.7E-01 AA470686.1	EST_HUMAN	Ine13a02.s1 NCI_CGAP_Co3 Homo sapiens cDNA clone IMAGE:881066 3' similar to gb:M17886 60S ACIDIC RIBOSOMAL PROTEIN P1 (HUMAN);
5820	1		12.31	1.7E-01		EST_HUMAN	ys02g06.s1 Soares fetal liver spleen 1NFLS Homo saplens cDNA clone IMAGE:213658 31
6112	1		2.15		1.7E-01 AF026552.3	Z	Mesocricetus auratus oviductin precursor (OVI) gene, complete cds

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SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acessian No.	Top Hit Database Source	Top Hit Descriptor
6320	16183	26344	7.96	1.7E-01	BE734179.1	EST_HUMAN	601569022F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3843984 5'
9288	16468	26858	1.2	1.7E-01	AF000573.1	NT	Homo sapiens homogentisate 1,2-dioxygenase gene, complete cds
6814	16693		7.03	1.7E-01	7708426 NT	TN	Homo saplens cleavage and polyadenylation specificity factor 3, 73kD subunit (CPSF3), mRNA
6814			7.03	1.7E-01	7706426 NT	FN FN	Homo sapiens cleavage and polyadenylation specificity factor 3, 73kD subunit (CPSF3), mRNA
7066	16943		2.48	1.7E-01	D00384.1	FX FX	Rat (SHR strain) SX1 gene
7531	17382	27593	7.38	1.7E-01	AP001508.1	N _T	Bacillus halodurans genomic DNA, section 2/14
7601	17452	27666	2.08	1.7E-01	U16288.1	NT	Human class IV alcohol dehydrogenase (ADH7), gene, exon 3
7875	17725		2.42	1.7E-01	AL 163284.2	NT	Homo sapiens chromosome 21 segment HS21C084
	l						Homo saplens solute carrier family 7 (cattonic anino acid transporter, y+ system), member 2 (SLC7A2),
7942	17792	28032	1.48	1.7E-01	11427203 NT		mRNA
7943	17793	28033	1.57	1.7E-01	AA627972.1	EST HUMAN	nq60e07.s1 NCI_CGAP_Co9 Homo sapiens cDNA clone IMAGE:1148292 3' similar to gb:L25081 TRANSFORMING PROTEIN RHOC (HUMAN);
8087	17958		9.13	1.7E-01	BE390835.1	EST HUMAN	601286547F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3613258 5'
8182	18068	28317	2.47	1.7E-01	AA814617.1	EST_HUMAN	of43a03.s1 NCI_CGAP_CNS1 Homo sapiens cDNA clone IMAGE:1426924 3'
8456	18329	28589	7.88	1.7E-01	7106300 NT	LY.	Mus musculus adenomatosis polyposis coli binding protein Eb1 (Eb1), mRNA
8456	18329	28590	7.88	1.7E-01	7106300 NT	TN	Mus musculus adenomatosis polyposis coli binding protein Eb1 (Eb1), mRNA
8949	18757		1.92	1.7E-01	P15272	SWISSPROT	AMP NUCLEOSIDASE
8012	18811	29107	4.38	1.7E-01	11418157 NT	NT	Homo saplens calcium channel, voltage-dependent, alpha 11 subunit (CACNA11), mRNA
9139	19666		1.5	1.7E-01	AL163278.2	NT	Homo sapiens chromosome 21 segment HS21Ct/78
9420	19513		1.28	1.7E-01	AI824404.1	EST HUMAN	t689g05.x1 NCI_CGAP_Ut1 Homo sapiens cDNA clone IMAGE:2274872 3′ similar to gb:M73779 RETINOIC ACID RECEPTOR ALPHA-1 (HUMAN);
9705	19253	25218	67.9	1.7E-01	U01317.1	LN	Human beta globin region on chromosome 11
120	10097	19917	1.88	1.6E-01	AF217532.1	NT	Homo sapiens mevalonate kinase gene, exon 6 and 7
664	12641	20416	1.53	1.6E-01	R31497.1	EST_HUMAN	yh75f12.r1 Soares placenta Nb2HP Homo saplen's cDNA clone IMAGE:135599 5'
1505	11409		4.05	1.6E-01	AF298117.1		Homo saplans homeobox protein OTX2 gene, coniplete cds
1882	11778	21653	2.4	1.6E-01	P22063	ISSPROT	AXONIN-1 PRECURSOR (AXONAL GLYCOPROTEIN TAG-1)
1941			1		U10334.1	TN	Crassostrea gigas RNA polymerase II largest subunit mRNA, partial cds
2335		22115	96.0	l		NT	H.sapiens mRNA for novel T-cell activation protein
2443	1		2.19			NT	Homo sapiens mRNA for KIAA1308 protein, partiel cds
2863	12791	22583	6.8	1.6E-01	AF185589.1	NT	Homo saplens cytochrome P450 3A4 (CYP3A4) gene, promoter region
2863	12791	22584	8.9	1.6E-01		NT	Homo saplens cytochrame P450 3A4 (CYP3A4) gene, promoter region
3581	13495	23285	1.31	1.6E-01	AJ003165.1	NT	Populus trichocarpa cv. Trichobel ABI3 gene
3581	1	23286	1.31			NT	Populus trichocarpa cv. Trichobel AB/3 gene
3919	13828		2.61	1.6E-01	AE004413.1	N	Vibrio cholerae chromosome II, section 70 of 93 of ithe complete chromosome

WO 01/57274

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					•		
Proba SEQ ID NO:	_ 8	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
4234	14132	23808	7.92	1.6E-01	AF179680.1	TN	Homo sapiens apelin gene, complete cds
4353	L		2.44	1.6E-01	AW968601.1	EST_HUMAN	EST380677 MAGE resequences, MAGJ Homo sapiens cDNA
4361	14257		4.01	1.6E-01	6753319 NT	TN	Mus musculus chaperonin subunit 3 (gamma) (Oct3), mRNA
4781	14665	24451	98.0	1.6E-01	Z28330.1	TN	S.cerevisiae chromosome XI reading frame ORF YKR105c
4781	14665	24452	0.86	1.6E-01	Z28330.1	NT	S.cerevisiae chromosome XI reading frame ORF YKR105c
4865	14745	24524	1.14	1.6E-01	AA088343.1	EST_HUMAN	284h09.st Stratagene colon (#937204) Homo saplens cDNA clone IMAGE:511361 3' similar to TR:E221955 E221955 38,855 BP SEGMENT OF CHROMOSOME XIV.;
4889	14769	24546	1.92	1.6E-01	AJ006356.1	Z	Lycopersicon esculentum Rsal fragment 2, satellite region
4889	14769	24547	1.92	1.6E-01	AJ006356.1	NT.	Lycopersicon esculentum Rsal fragment 2, satellite region
4958	14833	24601	1.09	1.6E-01	BE018707.1	EST_HUMAN	bb83h08.y1 NIH_MGC_10 Homo sapiens cDNA clone IMAGE:3049023 5' similar to gb:M61715 TRYPTOPHANYL-TRNA SYNTHETASE (HUMAN); gb:X69657 M.musculus (MOUSE);
6390	15309	25162	3.12	1.8È-01	AW197496.1	EST HUMAN	xm43f01.x1 NCI_CGAP_GC6 Homo saplens cDNA clone IMAGE:2686969 3' similar to TR:075984 O75984 HYPOTHETICAL 127,6 KD PROTEIN ;
000	j		6	4 05 04	7 0072077814	FOT LINAAN	xm43f01.xf NCI_CGAP_GC6 Home saplens cDNA clone IMAGE:2686969 3' similar to TR:075984 075964
200	١		3.12	10-10-1	AW 187490.1	ביין	Post and a second secon
5398	15317	25384	2.07	1.65-01	AF034/16.1	Z	Ratus norvegicus CCAA Fennancer piraing protein epsilon (cepte) gene, complete cas Arabidonete thellane DNA chromosome 4, conflictrament No. 84
5873	1	25899	2.24	1.65-01		NT.	Arabidopsis thaliana DNA chromosome 4, continuament No. 84
6157	┸	24858	3.7	1.6E-01	AW 291215.1	EST HUMAN	Ul-H-BI2-agi-b-06-0-Ul.s1 NCI_CGAP_Sub4 Homo saplens cDNA clone IMAGE:27244183'
6571	16428	26612	1.84	1.6E-01		EST_HUMAN	2822248.5prime NIH_MGC_7 Homo sapiens cDNA clone IMAGE:2822248 5'
6582	18472	28682	1.42	1.6E-01	L49349.1	NT	Gorilla gorilla androgen receptor gene, partial exon
7187	17064	27254	1.89	1.6E-01	249501.1	TN	S.cerevisiae chromosome X reading frame ORF YJR001w
7564	17415		1.7	1.6E-01	BF375171.1	EST_HUMAN	RC3-ST0200-041199-011-h01 ST0200 Homo sapiens cDNA
7565	17416	27631	1.91	1.6E-01	Z49501.1	NT	S.cerevisiae chromosome X reading frame ORF YJR001w
8049	17940	28190	2.71	1.6E-01	AW850853.1	EST_HUMAN	IL3-CT0220-111199-028-G01 CT0220 Homo sepiens cDNA
8364		28480	1.78	1.6E-01	014647	SWISSPROT	CHROMODOMAIN-HELICASE-DNA-BINDING PROTEIN 2 (CHD-2)
8364	18241	16487	1.78	1.6E-01	014647	SWISSPROT	CHROMODOMAIN-HELICASE-DNA-BINDING FROTEIN 2 (CHD-2)
8459	18332		7.6	1.6E-01	AF106064.1	NT	Plasmodium falciparum calcium-dependent protein kinase-3 (cdpk3) gene, complete cds
8713	18530	28814	10.07	1.6E-01	6671552 NT	NT	Mus musculus adaptor-related protein complex AF-1, beta 1 subunit (Ap1b1), mRNA
9001	18804	28082	2.69	1.6E-01	AW8771:	EST HUMAN	QV2-PT0010-160400-133-e08 PT0010 Homo sepiens cDNA
9025	Ц		2.17	1.6E-01	6679466 NT	NT	Mus musculus protein kinase, cGMP-dependent, type II (Prkg2), mRNA
9141	18896	28795	2.33	1.6E-01	AV719585.1	EST HUMAN	AV719585 GLC Homo sapiens cDNA clone GLCEMF07 5'
9565			6.33	1.6E-01		NT	Cucumis sativus KS mRNA for ent-kaurene synthase, complete cds
9727	19265		2.84	1.6E-01	AK024496.1	ᅜ	Homo saplens mRNA for FLJ00104 protein, partial cds

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Top Hit Descriptor	Fuchsia hybrid cultivar Qiu 94208 ribosomal protein S10 gene, partial cds; nuclear gene for milochondrial product	Rattus norvegicus chondroitin sulfate proteoglycan 5 (neuroglycan C) (Cspg5), mRNA	IL3-HT0619-040700-197-E05 HT0619 Homo saplens cDNA	IL3-HT0619-040700-197-E05 HT0619 Homo suplens cDNA	AV711696 DCA Homo sapiens cDNA clone DCAADH06 5'	Homo saplens chromosome 21 segment HS21C084	Opprinus carplo mRNA for EGGS22 myosin heavy chain, 3'UTR	Homo sapiens partial SLC22A2 gene for organic cation transporter (OCT2), exon 1	Rattus norvegicus insulin-responsive glucose transporter (GLUT4) gene, 5' end	xn39d11.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2696085 3'	Human gene for dihydrolipoamide succinyltransferase, complete cds (exon 1-15)	Human gene for dihydrolipoamide succinyltransferase, complete cds (exon 1-15)	Mus musculus MAP kinase kinase kinase 1 (Melkt1) mRNA, complete cds	UI-H-Bi3-akb-b-09-0-UI.s1 NCI_CGAP_Sub5 Homo sapiens cDNA clone IMAGE:2733641 3'	602083269F1 NIH_MGC_81 Homo sapiens cDNA clone IMAGE:4247537 5	Bos taurus factor V variant 2 (factor V) mRNA, complete cds	oo88d05.s1 NOLCGAP_GC4 Homo sapiens cDNA clone IMAGE:1571337 3' similar to gb:M11433 RETINOL-BINDING PROTEIN I, CELLULAR (HUMAN);	L.stagnalis mRNA for G protein-coupled receptor	L. stagnalis mRNA for G protein-caupled receptor	hh29f02,x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:2956539 3' similar to contains element MER16 repetitive element;	Mus musculus ICR/Swiss glyceraldehyde 3-phosphate dehydrogenase (Gapd-S) gene, complete cds	Homo saplens pyruvate dehydrogenase kinase, is senzyme 1 (PDK1), nuclear gene encoding mitochondrial	protein, mRNA	hj10f06.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2981411 3'	Populus trichocarpa cv. Trichobel ABI3 gene	Populus trichocarpa cv. Trichobel ABI3 gene	RC2-HT0149-191099-012-c09 HT0149 Homo saplens cDNA	B.napus mitochondrion DNA for ORF158	Homo sapiens chromosome 21 segment HS210034	602067192F1 NIH_MGC_57 Homo sapiens cDNA clone IMAGE:4066223 5'
Top Hit Database Source	TN	F	EST_HUMAN	EST_HUMAN	EST HUMAN	TN	. TN	NT	TN	EST_HUMAN	Z	NT	Į,	EST_HUMAN	EST_HUMAN	NT	EST HUMAN	N	N	EST_HUMAN	TN		NT	EST_HUMAN	IN	LΝ	EST_HUMAN	NT	NT	EST_HUMAN
Top Hit Acession No.	AF287344.1	9506522 NT	BE710087.1	BE710087.1	AV711696.1	AL163284.2	AJ009735.1	AJ251885.1		6.1	D26535.1	D26535.1	AF117340.1	AW44451.1	BF695381.1		AA935049.1	223104.1	223104.1	AW612237.1	U09964.1		7108358 NT	AW665983.1	AJ003165.1	AJ003165.1	AW366659.1	212628.1		BF687665.1
Most Similar (Top) Hit BLAST E Value		1.6E-01	1.5E-01	1.5E-01	1.5E-01	1.5E-01	1.5E-01	1.5E-01	1.5E-01	1.5E-01	1.5E-01	1.5E-01	1.5E-01	1.5E-01	1.5E-01		1.5E-01	1.5E-01	1.5E-01	1.5E-01	1.5E-01		1.5E-01	1.5E-01	1.5E-01	1.5E-01	1.5E-01	1.5E-01		1.5E-01
Expression Signal	2.47	1.27	1.87	1.87	2.28	1.4	0.88	1.87	1.82	1.36	2.81	2.81	1.49	1.54	1.12	0.89	4.22	78.0	78.0	98:0	1.34		185.26	2.35	8.0	9.0	0.98	0.97	8.36	1.34
ORF SEQ ID NO:			20030	20031		20534		20838		20953	21010		21238	21636	22435		23034			23087	23394			23490	90582	23507	23649	23687		24301
Exen SEQ ID NO:	19319	19328	10214	10214	12640	10697	10992	10997	11013	11108	11161	11161	11371	11762	12544	l.	13229	13242	13242	13298	13610			13703	13718	13718	13871	13912		14511
Probe SEQ ID NO:	8807	9827	248	248	572	766	1076	1081	1087	1198	1254	1254	1466	1866	2879	2999	3308	3322	3322	3380	3696		3706	3791	3808	3806	3984	4008	4091	4623

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Table 4
Single Exon Probes Expressed in Heart

Probe SEQ ID NO:	Ecan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
4645	12844		2.18	1.6E-01	BF695381.1	EST_HUMAN	602083269F1 NIH_MGC_81 Hamo sapiens cDNA clane IMAGE:4247537 5'
4680	14568	24361	1.08	1.5E-01	BE173796.1	EST_HUMAN	CM0-HT0565-280200-245-b10 HT0565 Homo saplens cDNA
4680	14566	24362	1.08	1.5E-01	BE173798.1	EST_HUMAN	CM0-HT0565-280200-245-b10 HT0565 Homo saplens cDNA
4929	14808		1.3	1.6E-01	AL161560.2	LZ	Arabidopsis thaliana DNA chromosome 4, contigifragment No. 60
6072	1			1.5E-01	AF003105.1	LΝ	Arabidopsis thaliana AP2 domain containing protein RAP2.12 mRNA, partial cds
5216	15139	24833	}	1.5E-01	P07996	SWISSPROT	THROMBOSPONDIN 1 PRECURSOR
2000	76700		60 9	70 27 7	044400	TOGGGGIA	SEX HORMONE-BINDING GLOBULIN PRECURSOR (SHBG) (SEX STEROID-BINDING PROTEIN) (SBD) (TEXTIS, SPECIAL ANDROGEN, BINDING PROTEIN) (ARD)
0070	- }		20.0	10-UC-1		DA PENA	
25 25 26 26 26 26 26 26 26 26 26 26 26 26 26	J			1.5E-01	4	EST_HUMAN	L3-C (0219-160200-064-F10 C (0219 Hamo saplens cUNA
6424				1.5E-01	U65018.1	NT	Mus musculus transforming growth factor alpha (TGFa) mRNA, complete cds
5424			22.9	1.5E-01	U65016.1	NT	Mus musculus transforming growth factor alpha (TGFa) mRNA, complete cds
5652	15564		1.9	1.5E-01	6753659 NT	LN	Mus musculus DNA methyltransferase 2 (Dnmt2), mRNA
5652			1.9		6753659 NT	NT	Mus musculus DNA methyliransferase 2 (Dnmt2), mRNA
5684	_		1.87	1.5E-01	AJ276505.1	Ę	Mus musculus genomic fragment, 279 Kb, chroniosome 7
5760	15668	25774	2.44	1.5E-01	BE727658.1	T_HUMAN	601564322F1 NIH_MGC_20 Homo sapiens cDNA clone IMAGE:3833981 5'
5785	15691		1.66	1.5E-01	45063BB NT	FA.	Homo saplens RAD54 (S.cerevislæe)-like (RAD54L) mRNA
8008)		1 70	4 10 10	A 5424007 1	FN	inflierza B virus (B.N.anchand/480/94.) NB profein cene complete cris: and neuraminidase cene nartial cris
3	5				Ī		Annual Control of the
5917	-)			1.5E-01	AE001039.1		Archaeoglobus fulgidus section os of 1/2 of the complete genome
5935	15840	25963	5.13	1.5E-01	11417236 NT	-N	Homo sapiens chromosome 5 open reading frame 3 (C5ORF3), mRNA
5942	15847	25971	1.81	1.5E-01	P48508	SWISSPROT	GLUTAMATECYSTEINE LIGASE REGULATORY SUBUNIT (GAMMA-GLUTAMYLCYSTEINE SYNTHETASE) (GAMMA-ECS) (GCS LIGHT CHAIN)
5972				1.5E-01	Q28462	SWISSPROT	AMELOGENIN
9031	15935	L	1.4	1.5E-0	P30143	SWISSPROT	HYPOTHETICAL 51.7 KD PROTEIN IN THRC-TALB INTERGENIC REGION (ORF8)
6168	15123		5.63	1.5E-01	AW970295.1	EST_HUMAN	EST382376 MAGE resequences, MAGK Hamo sapiens cDNA
6284			1.77	1.5E-01	AF210842.1	NT	Homo sapiens HARP (HARP) gene, excn 17 and complete cds
6374			1.88	1.5E-01	AI973157.1	EST_HUMAN	wr52c08.x1 NCI_CGAP_Utf Homo sapiens cDNA clone IMAGE:2491310 3'
6481		26507		1.5E-01		EST_HUMAN	UI-HF-BNO-akk-d-05-0-UI.r1 NIH_MGC_50 Hamp saplens cDNA clone IMAGE:3077409 5'
6481	16340		1.68	1.5E-01	AW500611.1	EST_HUMAN	UI-HF-BN0-akk-d-05-0-UI.r1 NIH MGC 50 Homo sapiens cDNA clone IMAGE:3077409 5
6788	16665	28859	1.22	1.5E-01	AA970317.1	EST HUMAN	0085g12.s1 NCI_CGAP_Kid5 Homo saplens cDNA clone IMAGE:1573030 3' simiter to gb:M26062 INTERLEUKIN-2 RECEPTOR BETA CHAIN PRECURSOR (HUMAN);
6895	16774		11.77	1.5E-01	C16800.1	EST HUMAN	C16800 Clontech human acrta polyA+ mRNA (#6572) Homo saplens cDNA clone GEN-529H09 5'
6912	1 1	26983		1.5E-01		TN	Pangasianodon gigas growth hormone (GH) mRNA, complete cds
9669			1.44	1.5E-0	D84476.1	NT	Homo sapiens mRNA for ASK1, complete cds

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Probe SEQ ID NO:	Exon SEQ:(D NO:	ORF SEQ ID NO:	Expression Signel	Most Similar (Top) Hit BLAST E Vatue	Top Hit Acesslon No.	Top Hit Database Source	Top Hit Descriptor
7130	17007	27200	1.71	1.5E-01	4501972 NT	Ŋ	Homo sapiens adaptor-related protein complex 1, beta 1 subunit (ADTB1), mRNA
7,005	474.40		07.0		4	144741111111111111111111111111111111111	2a59e08.s1 Scares fetal liver spleen 1NFLS Homo saplens cDNA clone IMAGE:296866 3' similar to
2007	7 27	57.333	2,40		11/4/20.1	NAMOR I CO	PINCE CATALON CALLES PROBINITION OF THE PROPINE TO
8	1/182		2.98		AV754819.1	EST HUMAN	AV754819 IP Homo saplens cDNA clone I PAAHB12 5
7438 88	16451		6.6		U00455.1	뒫	Acipenser transmontano vitellogenin mRNA, partial cds
7708	17556		7.02		AF007570.1	Į.	Aplysia californica carboxypeptidase D mRNA, complete cds
7708	17556		7.02		AF007570.1	NT	Aplysia californica carboxypeptidase D mRNA, complete cds
7860	17710	27955	2.86		1.5E-01 X98852.1	NT	P. Ieniusculus mRNA for Integrin beta subunit
7908	17758	27998	2.45	1.6E-01 A	AI814046.1	EST_HUMAN	wk33h12x1 NCI_CGAP_Pr22 Homo sapiens cDNA clone IMAGE:2419175 3' similar to gb:M27508 BETA
7908	17758	27999	2.45		1.5E-01 AI814046.1	EST HUMAN	wkS3h12.x1 NCI_CGAP_Pr22 Homo septens cDNA clone IMAGE:2419175 3' similar to gb:M27508 BETA GALACTOSIDASE-RELATED PROTEIN PRECURSOR (HUMAN):
7939	17789	28031	1.54		U40932.1	Ę	Danio reno transcription factor PaxBb (PaxB) mRNA, complete cds
8008	17859	28103	1.35		ıσ	E	Claviceps purpurea ps1 gene
8009	17859		1.35		1.5E-01 AJ011964.1	FZ	Claviceps purpurea ps1 gene
8199	18084	28334	5.15	, ,	1.5E-01 AL163280.2	NT	Homo sapiens chromosome 21 segment HS21C080
8199	18084	28335	5.15		1.5E-01 AL163280.2	NT	Homo saplens chromosome 21 segment HS21C080
8342	18219		1.74		1.5E-01 AB042975.1	NT	Sus scrofa CYP51 gene for lanosterol 14 alpha-demethylase, exon 1
8425	18299		1.73		1.5E-01 AW841915.1	EST_HUMAN	IL5-CN0024-030300-025-D04 CN0024 Homo sapiens cDNA
8506	16236	26396	2.17		1.6E-01 AI973157.1	EST HUMAN	wr52c08x1 NCL_CGAP_Ut1 Homo sepiens cDNA clone IMAGE:2491310 3'
6606	19547		20.02		BF700582.1	EST_HUMAN	602128753F1 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:4285549 5'
9531	19565		4.14		R83077.1	EST_HUMAN	yp87e04.r1 Soares fetal liver spicen 1NFLS Homo sapiens cDNA clone IMAGE:194430 5'
9621	19588		2.14	١	1.5E-01 AV741272.1	EST_HUMAN	AV741272 CB Homo sapiens cDNA clone CBDA GD04 5'
9728	19497		3.99	•	1.5E-01 AL139074.2	TN	Campylobacter jejuni NCTC11168 complete genome; segment 1/6
9932	19402	25179	3.01	1.5E-01	1.5E-01 AJ276242.1	IN	Sus scrofa mRNA for sodium iodide symporter
9972	19432		1.58		1.5E-01 AF020346.1	TN	Rattus norvegicus pyridoxal kinase mRNA, complète cds
296	10280		1.96		1.4E-01 AF009663.1	Ł	Homo sapiens T cell receptor beta locus, TCRBViSS5P to TCRBV21S2A2 region
892	10818		2.57	1.4E-01	D78638.1	TN	Xenopus laevis mRNA for DNA (cytosine-5-)-methyltransferase, complete cds
1238	11145		1.62		T91864.1	EST_HUMAN	yd54c01.s1 Soares fetal liver spleen 1NFLS Home sapiens cDNA clone IMAGE:112032 3'
1714	11615		1.43		TN 0868289	TN	Mus musculus growth differentiation factor 5 (Gdf3), mRNA
1717	11618	21487	1.53	1.4E-01	1.4E-01 AE001710.1	LN	Thermotoga maritima section 22 of 136 of the complete genome
1863	11769		0.94		1.4E-01 AW135741.1	EST_HUMAN	UI-H-BI1-acf-a-09-0-UI.s1 NCI_CGAP_Sub3 Horno sapiens cDNA clone IMAGE:2714009 3'
1942	11837		10.35		1.4E-01 AA720815.1	EST_HUMAN	ny72d07.s1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:1283821 3'
2426	12303	22189	0.97		P30706	SWISSPROT	GLYCEROL-3-PHOSPHATE ACYLTRANSFERASE PRECURSOR (GPAT)

Page 76 of 413 Table 4 Single Exon Probes Expressed in Heart

		### 17401.X1 NOI_CGAP_UZ Homo saplens cDNA clone IMAGE:2441665 3' ### 17401.X1 NOI_CGAP_UZ Homo saplens cDNA clone IMAGE:341467 5' ### 17501.Soares infant brain 1NIB Homo saplens cDNA clone IMAGE:3273570 3' ### 17501.Soares infant brain 1NIB Homo saplens cDNA clone IMAGE:2273570 3' ### 17501.Boares infant brain 1NIB Homo saplens cDNA clone IMAGE:2273570 3' ### 17501.Boares infant brain 1NIB Homo saplens cDNA clone IMAGE:3273570 3' ### 17501.Boares fetal_liver_spleen_1NIPLS_51 Homo saplens cDNA clone IMAGE:44467 5' ### 17501.Boares infantsacion 22 of 136 of the complete genome ### 17501.Boares infantsacion 22 of 136 of the complete cDNA ### 17501.Boares infantsacion 22 legital_liver_spleen_1NIPLS_51 Homo saplens cDNA ### 17501.Boares infantsacion 22 legital_liver_spleen_1NIPLS_51 Homo saplens cDNA ### 17501.Boares infantsacion 22 legital_liver_spleen infantsacion 22 legital_liver_spleen infantsacion 22 legital_liver_spleen infantsacion 22 legital_liver_spleen infantsacion 22 legital_liver_spleen infantsacion saplens cDNA clone IMAGE:113838 3' ### 17501.Boares infantsacion 22 legital_liver_spleen infantsacion 22 legital_liver_spleen infantsacion 23 legital_liver_spleen infantsacion 24 legital_liver_spleen infantsacion infa	THUMAN THUMAN THUMAN THUMAN THUMAN THUMAN THUMAN THUMAN THUMAN THUMAN THUMAN THUMAN	AB933496.1 R69232.1 AR692932.1 AR692934.1 AR692094.1 AR692094.1 AR692094.1 AR692094.1 AR692094.1 AR692094.1 AR692094.1 AR692094.1 AR692094.1 AR692094.1 AR692094.1 AR692094.1 AR692094.1 AR6920947.1	1.4E-01 1.4E-0				2762 3823 3823 4083 4083 4144 4144 41144 4313 5267 5267 5267 5267 5267 5267 5268 5268 5268 5268 5268 5268 7368 7368 7368 7368 7368 7368 7368 73
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Exon SEQ ID NO: ORF SEQ Signal Signal Signal Expression (Top) Hit BLASTE Value Top Hit Acession No. Top Hit Acession No. Top Hit Acession Source Value Top Hit Acession No. Top Hit Source 12624 22517 3.07 1.4E-01 Al933498.1 EST HUMAN 13736 23524 1.4E-01 R59232.1 EST HUMAN	Exan ORF SEQ ID NO: Expression Signal NO: Top Hit Acession (Top) Hit Acession ID No: Top Hit Acession No: Top Hit Acession Database Source Value 12624 22517 3.07 1.4E-01 Al933496.1 EST HUMAN 13736 23524 1 -4E-01 R99232.1 EST HUMAN	Solates intent ordin Tinib Hand saplens cone IMAGE:41407.5				1			3823
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Most Similar	Most Similar	Top Hit Descriptor		No.	BLAST E Value	Signal		SEQ ID	Ωö
		Top Hit Descriptor		Top Hit Acession No.	(Top) Hit BLAST E Value	Expression Signal		Exan SEQ ID	eg c ö

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Table 4
Single Exon Probes Expressed in Heart

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Тор Hit Descriptor
8764	17913	28158	2.23	1.4E-01	U28760.1	LN	Borrelia burgdorferi glyceraldehyde-3-phosphata dehydrogenase (GAPDH), phosphoglycerate kinase (PGK), triosephosphate isomerase (TPI) genes, complete cds
8813	18626		3.02	1.4E-01	X52102.1	NT	M.musculus p16K gene for 16 kDa protein
9365	19517	25138	1.48	1.4E-01	AB000890.1	NT	Ephydatia fluviatilis mRNA for alddase, partial cds
9413	19065	25277	2.32	1.4E-01	X74773.1	TN	P.salina plastid gene secY
9427	19073		1.89	1.4E-01		TN	Rettus norvegicus desmin (Des), mRNA
9470	19734		1.82	1.4E-01	BE513802.1	EST_HUMAN	601315638F1 NIH_MGC_8 Homo sapiens cDNA clone IMAGE:3634329 5'
0580	10158		3 04	10 37 1	AE083224 4	F	Fugu rubribes putative neurotransmitter receptors, YDR140w homolog, and glycinamide ribonucleotide transformulase (CART) names complete ets
9573	1		2.29	1.4E-01	_	ΙZ	Synechocystis sp. PCC6803 complete genome, 29/27, 2868767-3002965
9646	19754		3.28	1.4E-01		SWISSPROT	TYROSINE-PROTEIN KINASE TRANSFORMING PROTEIN ABL
9762	19282		1.41	1.4E-01		EST_HUMAN	601465575F1 NIH_MGC_67 Homo saplens cDNA clone IMAGE:3868795 5'
9831	19332		1.42	1.4E-01	11425031		Homo sapiens ephrin-B3 (EFNB3), mRNA
9850	19566		3.41	1.4E-01	D82983.1	N L	Mus musculus mRNA for prolidase, complete cds
9356	19398	ŀ	1.7	1.4E-01	AW377998.1	EST_HUMAN	MRo-HT0208-221299-204-c08 HT0208 Homo saplens cDNA
319	10281	20098	2.69	1.3E-01	4758467 NT	N	Homo sapiens G protein-coupled receptor 50 (GPR50) mRNA
319	10281	20099	2.69	1.3E-01	4758467 NT	TN	Homo saplens G protein-coupled receptor 50 (GPR50) mRNA
518	10460		1.86	1.3E-01	AB013139.1	NT	Homo sapiens gene for NBS1, complete cds
620	10557	20369	0.89	1.3E-01	AJ277606.1	TN	Human calicivirus HU/NLV/Girlington/93/UK RNA, for capsid protein (ORF2), strain HU/NLV/Girlington/93/UK
620	10557	20370	0.89	1.3E-01	AJ277608.1	LN.	Human calicivirus HU/NLV/Girlington/93/UK RNA for capsid protein (ORF2), strain HU/NLV/Girlington/93/UK
826	10763	20603	1.09	1.3E-01	X53330.1	Z	P.dumerilii histone gene cluster for core histones H2A, H2B, H3 and H4
876	10802	20852	1.83	1.3E-01	AF139518.1	NT	Rattus norvegicus A-kinase anchor protein mRNA, complete cds
1010	10928	20771	1.55	1.3E-01	AL117078.1	NT	Botrytis cinerea strain T4 cDNA library under conditions of nitrogen deprivation
1111	11026		2.23	1.3E-01	AL115265.1	NT	Botrytis cinerea strain T4 cDNA library under conditions of nifrogen deprivation
1197	11107	20922	1.07	1.3E-01	AV712467.1	EST_HUMAN	AV712487 DCA Homo sapiens cDNA clone DCAAFF05 5'
1426	11331		1.36	1.3E-01		NT -	Homo sapiens adapter protein CMS mRNA, complete cds
1916	11811	21689	2.56	1.3E-01	AL117078.1	ΝT	Botrytis cinerea strain T4 cDNA library under conditions of nitrogen deprivation
2124	12012	·	1.29	1.3E-01	AJ243578.1	TN	Rhodopseudomonas acidophila pucBS, pucAS, pucAS, pucAS, pucB7, pucA7, pucB8, pucA8 and pucC genes and ORF151
2245	12129		1.17	1.3E-01		EST_HUMAN	RC4-ST0173-191099-032-d12 ST0173 Homo saplens cDNA
2329	12210		2.89			NT	Archaeoglobus fulgidus section 91 of 172 of the complete genome
2542	12416	22306	3.49	1.3E-01	M86918.1	<u>N</u>	Carassius auratus keratin type I mRNA, complete cds

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	Top Hit Descriptor	Bowne branched chain alpha-keto acid dihydrolipcy/ transacylase mRNA, complete cds	Pyrococcus harikoshii OT3 genamic DNA, 1-287000 nt. position (1/7)	Pyrococcus harikoshii OT3 genomic DNA, 1-287000 nt. position (1/7)	Homo sapiens DD4 gene for dihydrodiol dehydrogenase 4 [AKR 1C4], exon 2	Pyrococcus harikoshii OT3 genomic DNA, 1-287000 nt. position (1/7)	Pyrococcus harlkoshil OT3 genomic DNA, 1-287000 nt. position (1/7)	Rattus norvegicus Fibrinogen, gamma polypeptide (Fgg), mRNA	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 77	Bactertophage SPBc2 complete genome	QV3-DT0018-081299-036-a03 DT0018 Homo sapiens cDNA	Schistosoma mansoni fructose bisphosphate aldolase mRNA, complete cds	xv23f10.x1 Soares_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:2813995 3'	AV752279 NPD Homo sapiens aDNA clane NPDAZE02 5'	AV752279 NPD Homo sapiens cDNA clone NPDAZE02 5	Homo sapiens chromosome 21 segment HS21C080	601126096F1 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:2990063 5	802154306F1 NIH_MGC_83 Homo sapiens cDNA done IMAGE:4285544 5	601510347F1 NIH_MGC_71 Homo saplens cDNA clone IMAGE:3911987 5	AU136819 PLACE1 Homo saplens cDNA clone FLACE1004693 5'	Homo sapiens chromosome 21 segment HS210034	Homo sapiens chromosome 21 segment HS21C084	602154401F1 NIH_MGC_83 Homo saplens cDNA clone IMAGE:4295305 57	602154401F1 NIH_MGC_83 Homo sapiens cDNA clone IMAGE:4295305 5	QV0-UM0093-100400-189-a06 UM0093 Homo sepiens cDNA	Hepatitis C virus 68_CL10 genome polyprotein gene, partial cds	Schizosaccharomyces pombe gene for Alp41, complete cds	Clacchus intron 4 of visual pigment gene (red allele)	yr33d02.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:207075 5'	Homo sapiens PRO0611 protein (PRO0611), mRNA	602187015T1 NIH_MGC_49 Homo sapiens cDNA clone IMAGE:4299074 3'	S.cerevisiae chromosome IV reading frame ORF YDL054c	Homo sapiens core histone macroH2A2.2 (MACROH2A2), mRNA	602187015T1 NIH_MGC_49 Homo sapiens cDNA clone IMAGE:4299074 3'	Oryctolagus cuniculus H+,K+-ATPase alpha 2c subunit mRNA, complete cds
	Top Hit Database Source	ŊŢ	NT	ΤN	F	NT	TN	IN	TN	NT	EST_HUMAN	TN	EST_HUMAN	EST_HUMAN	EST_HUMAN	Ν	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	INT	N	EST_HUMAN	EST_HUMAN	EST_HUMAN	INT	TN	TN	EST_HUMAN	N.	EST_HUMAN	TN	LN.	EST_HUMAN	NT
	Top Hit Acession No.	M21572.1	AP000001.1	AP000001.1	AB032159.1	AP000001.1	AP000001.1	6978840 NT	AL161581.2	AF020713.1	AW364341.1	AF026805.1	AW273741.1	AV752279.1	AV752279.1	AL163280.2	BE272339.1	BF679654.1	BE884017.1	AU136619.1	AL163284.2	AL163284.2	BF679819.1	BF679819.1	AW804417.1	AF056880.1	AB031326.1	X88891.1	H48684.1	11423294 NT	BF690522.1	Z74102.1	8923919 NT	BF690522.1	AF023129.1
	Most Similar (Top) Hit BLAST E Value	1.3E-01	1.3E-01	1.3E-01	1.3E-01	1.3E-01	1.3E-01	1.3E-01	1.3E-01	1.3E-01	1.3E-01		1.3E-01			1.3E-01		_			1.3E-01		1.3E-01	1.3E-01		1.3E-01	1.3E-01		1.3E-01	1.3E-01	1.3E-01	1.3E-01	1.3E-01		
	Expression Signal	0.95	1.18	1.18	0.78	0.86	0.86	0.85	1.7	1.15	3.44	1.80	16.36	0.85	0.85	1.65	2.16	0.81	3.17	98.0	1.21	121	6.0	6.0	2.51	1.79	13.21	2.04	2	1.34	1.28	4.54	4.14	1.27	4.45
	ORF SEQ ID NO:	23122	23362		23368		23363	23431				23758	23770		23871		24118				24718	24719	24805	24806	25013			25983		26723					Ш
	Exon SEQ ID NO:	13321	13575	13575	13581	13575	13575	13648	13816	13948	13968	13977	13993	14092	14092	14116	14328	14418	15075	14783	14944	14944	15039	15039	15213	15348	16810	15861	16169	16529	16548	16737	18765	16838	17261
	Probe SEQ ID NO:	3404	3861	3661	3667	3714	3714	3734	3908	4046	4068	4075	4083	4182	4192	4218	4433	4525	4771	4903	5074	5074	5173	5173	5292	5428	6904	5956	6305	6848	9999	6858	9889	0960	7452

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Tcp Hit Descriptor
8029	17921		2.88	1.3E-01	BF330999.1	EST_HUMAN	MR4-BT0358-130700-010-h08 BT0358 Homo saplens cDNA
8444	18318	28577	1.83	1.3E-01	AF119117.1	NT	Homo sapiens dopamine transporter (SLC6A3) gene, complete cds
8578	18444		5.13	1.3E-01	6871745 NT	۲	Mus musculus cofilin 2, muscle (Cf2), mRNA
8873			3.72		BE279449.1	EST_HUMAN	601158052F1 NIH_MGC_21 Homo sapiens cDi\A clone IMAGE:3504804 5'
9261	18968	25320	1.97	1.3E-01	BE618346.1	EST HUMAN	601462741F1 NIH_MGC_67 Homo septens cDNA clone IMAGE:3866003 5'
6686	18054		3.18	1.3E-01	AJ242790.1	TN	Gallus gallus scyc1 gene for lymphotactin, exons 1-3
9757	19279		1.56	1.3E-01	AB026829.1	TN	Ephydatia fluviatilis mRNA for sALK-6, complete,cds
9784	19297		1.32	1.3E-01	AW001114.1	EST_HUMAN	wu24d09.x1 Soares_Dieckgraafe_colon_NHCD Homo sapiens cDNA clone IMAGE:2520977 3' similar to TR:060287 060287 KIAA0539 PROTEIN.;
9945	19414		1.26	1.3E-01	BF571764.1	EST_HUMAN	602078440F1 NIH_MGC_62 Homo saplens cDNA clone IMAGE:4253049 5'
378	10362	20185	7.21	1.2E-01	AI421744.1	EST_HUMAN	f39b02.x1 NCI_CGAP_Bm23 Homo sapiens cDNA clone IMAGE:2098639 3' similar to gb:U05760_ma1 ANNEXIN V (HUMAN);
418	9889		1.55	1.2E-01	U66912.1	TN	Dictyostellum discoldeum ORF DG1016 gene, perfial cds
535	10476		2.63	1.2E-01	AF039442.1	TN	Homo sapiens colon cancer antigen NY-CO-45 mRNA, partial cds
1355	11261	21117	2.78	1.2E-01	AU149146.1	EST_HUMAN	AU149146 NT2RM4 Homo sepiens cDNA clone NT2RM4001691 3'
1355	11261	21118	2.78	1.2E-01	AU149146.1	EST_HUMAN	AU149146 NT2RM4 Homo sapiens cDNA clone NT2RM4001691 3'
1361	11267		3.94	1.2E-01	AV735249.1	EST_HUMAN	AV735249 cdA Homo saplens cDNA clone cdAAJB11 5'
1490	11395		1.13	1.2E-01	AA897474.1	EST_HUMAN	al48e09.s1 Soares_NPL_T_GBC_S1 Homo septions cDNA clone IMAGE:1460584.3' simitar to TR:Q16671 Q16671 ANTI-MULLERIAN HORMONE TYPE II RECEPTOR PRECURSOR.;
1613	11517	21377	1.17	1.2E-01	014934	SWISSPROT	NUCLEAR FACTOR OF ACTIVATED T-CELLS! CYTOPLASMIC 4 (T CELL TRANSCRIPTION FACTOR NFAT3) (NF-ATC4) (NF-AT3)
1831	<u> </u>		2.62	1.2E-01	AI285402.1	EST_HUMAN	qt69f09.x1 NCI_CGAP_Eso2 Home sapiens cDNA clone IMAGE:1960553 3'
1730	11631		29.48	1.2E-01	X89211.1	TN	H.sepiens DNA for endogenous retroviral like element
1877	11773	i	1.43	1.2E-01	AW 449368.1	EST_HUMAN	UFH-BI3-aki-e-10-0-UI.s1 NCI_CGAP_Sub5 Homo sapiens cDNA clone IMAGE:2734564 3'
2134	12022	21919	2.1	1.2E-01	BF248490.1	EST_HUMAN	60182/1567F1 NIH_MGC_62 Homo saplens cDNA clone IMAGE:4046224 5'
2240	12124	22022	1.01	1.2E-01		TN	Homo sapiens chromosome 21 segment HS21C013
2546	12420		2.02	1.2E-01	AW996556.1	EST HUMAN	QV3-BN0046-220300-129-f10 BN0046 Homo sapiens cDNA
							ts18g07.x1 NC_CGAP_Pan1 Homo sapiens cDNA clone IMAGE:2228988 3' similar to TR:Q14048 Q14048 COLLAGEN VI ALPHA-2 ALTERNATIVE C-TERMINAL DOMAIN. [1] ;contains element PTR5 repetitive
2697	12561	22451	0.86	1.2E-01	AI623388.1	EST_HUMAN	element;
2812	12741	22537	1.5	1.2E-01	U18018.1	NT	Human E1A enhancer binding protein (E1A-F) mRNA, partial cds
2872	12799	22694	1.96	1.2E-01	AI720470.1	EST_HUMAN	as80c09.x1 Barstead colon HPLRB7 Homo sepieris cDNA clone IMAGE:2335024 3' similar to gb:L05095 60S RIBOSOMAL PROTEIN L30 (HUMAN);
2904	12831	22628	2.89	1.2E-01	M16364.1	NT	Human creatine kinase-B mRNA, complete cds

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_		_	_			_		_	_	_			_	_	_	_	_	_	_		_				_	-	-	_		_	_
	Top Hit Descriptor	Wheat mRNA for a group 3 late embryogenesis abundant protein (LEA)	QV1-BT0259-261099-021-d05 BT0259 Homo saplens cDNA	Methanococcus jannaschli section 142 of 150 of the complete genome	Bacilius subtilis complete genome (section 15 of 21); from 2795131 to 3013540	Wheat mRNA for a group 3 late embryogenesis abundant protein (LEA)	Wheat mRNA for a group 3 late embryogenesis abundant protein (LEA)	Bacillus subtilis complete genome (section 15 of 21): from 2795131 to 3013540	601810786R1 NIH_MGC_46 Homo saplens cDNA clone IMAGE:4053668 3'	P.clarkii mRNA; repeat region (ID 2MRT7)	P.clarkil mRNA; repeat region (ID 2MRT7)	HEMOLYSIN PRECURSOR	HYPOTHETICAL 52.4 KD PROTEIN C12B10.08C IN CHROMOSOME I	HYPOTHETICAL 52.4 KD PROTEIN C12B10.08C IN CHROMOSOME I	UI-HF-BK0-aah-d-01-0-UI.r1 NIH_MGC_38 Home sapiens cDNA clone IMAGE:3053817 5'	2c08d02.r1 Soares_parathyroid_tumor_NbHPA Homo sapiens cDNA clone IMAGE:321699 5'	Homo sapiens gene encoding plakophilin (exons 1-13)	801493518F1 NIH_MGC_70 Homo saplens cDNA done IMAGE:3895613 5'	IL0-CT0031-221099-113-604 CT0031 Homo sapiens cDNA	Mouse galactosyltransferase mRNA, complete cds	PM3-BN0137-290300-002-f09 BN0137 Homo saplens cDNA	wc99g03.x1 NCI_CGAP_Co3 Homo sapiens cDNA clone IMAGE:2326804 3' similar to SW:GST2_HUMAN Q99735 MICROSOMAL GLUTATHIONE S-TRANSFERASE II;	xc49d07.x1 NCI_CGAP_Eso2 Homo sapiens cDNA clone IMAGE:2587597 3' similar to gb:M13452 LAMIN A (HUMAN);	Staphylococcus aureus plasmid pSK23 putative recombinase Sin (sin) gene, partial cds; and transcriptional permitters Care Complete Cds.	Haemophilus influenzae Rd section 29 of 163 of the complete genome	S.cerevislae HXT6 gene	AV710857 Cu Homo sapiens cDNA clone CuAAKE08 5	Yeast MPT5 gene for suppressor protein, complete cds	601655578R1 NIH_MGC_65 Homo septens cDNA clone IMAGE:3846283 3'	801900763F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:4130103 5'	Homo sapiens dynein intermediate chain DNAI1 (DNAI1) gene, exon 17
	Top Hit Database Source	NT	EST_HUMAN	ΙN	NT	NT	TN	TN	EST_HUMAN	TN	NT	SWISSPROT	SWISSPROT	SWISSPROT	EST_HUMAN	EST_HUMAN	TN	EST_HUMAN	EST_HUMAN	NT	EST_HUMAN	EST_HUMAN	EST HUMAN	FIA	LZ	LN N	EST HUMAN	N.	EST_HUMAN	EST_HUMAN	NT.
	Top Hit Aœsslon No.	X56882.1	AW370668.1	U67600.1	299118.1	X56882.1	X56882.1	299118.1	BF128551.1	Z54255.1	Z54255.1	P16466	Q10441	Q10441	AW401836.1	W33035.1	Z98266.1	BE620945.1	AW845275.1	M26925.1	BE007072.1	AI913753.1	AW083652.1	AE069777 4	U32714.1	X77961.1	AV710857.1	D26184.1	BE962324.2	BF314481.1	AF190493.1
	Most Similar (Top) Hit BLAST E Value	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	, C	1.2E-01		1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01
	Expression Signal	0.98	2	76.0	0.79	1.14	1.14	1.2	88.0	2.2	2.2	1.04	1.47	1.47	2.47	2.63	1.9	1.69	2.19	1.56	1.21	2.46	9.72	0	227	10	1.51	2.85	3.35	1.93	2.6
	ORF SEQ ID NO:		22823				23199			23766	23767		24807	24808		24947	25021	25778	25824	25858		26715				27546					28715
	Exan SEQ ID NO:	12901	13117	13143		13393	13393	13350	13617	13990	13990	14886	15040	15040	15062	15174	15218	15689	15711	15745	16487	16522	16772	6797	L		17597		18291	IJ	18447
	Probe SEQ ID NO:	2974	3182	3219	3433	3477	3477	3554	3704	4090	4090	5012	5174	5174	5199	6251	5297	5762	9089	5839	6607	6842	6883	g	7043	7521	7747	8260	8417	8487	8579

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Top Hit Descriptor	Rabbit glycogen-associated protein phosphatase regulatory subunit (RG1) mRNA, complete cds	AV658033 GLC Homo sapiens cDNA clone GLCFIB12 3'	Homo saplens Xq pseudoautosomal region; segment 2/2	MACROPHAGE-STIMULATING PROTEIN RECEPTOR PRECURSOR (MSP RECEPTOR) (P185-RON) (CDW136) (CD138 ANTIGEN)	Homo sapiens colon cancer antigen NY-CO-45 mIRNA, partial cds	R.norvegicus NF68 gene for 68kDa neurofilament	QV4-BT0234-111199-031-g10 BT0234 Hamo saplens cDNA	qn20g05.x1 NCL_CGAP_Lu5 Homo sapiens cDNA done IMAGE:1898840 3'	Xenopus laevis integrin alpha 3 subunit mRNA, partial cds	CYCLIN T	601900763F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:4130103 5	th18d08.x1 NCI_CGAP_Brn25 Homo saplens cDNA clone IMAGE:2167983 3'	nm08g11.s1 NCI_CGAP_Co10 Homo sapiens cDNA clone IMAGE:1059820 3' similar to gb:X06985_ma1	HEME OXYGENASE 1 (HUMAN);	602128847F1 NIH_MGC_66 Homo sapiens cDNA clone IMAGE:4286771 5	Arabidopsis thallana DNA chromosome 4, contig fragment No. 60	EST384142 MAGE resequences, MAGL Hamo saplens.cDNA	Synechocystis sp. PCC6803 complete genome, 23/27, 2868767-3002965	AU140363 PLACE2 Homo sapiens cDNA clone PLACE2000403 5'	Mus musculus pre T-cell antigen receptor alpha (Picra), mRNA	Rattus norvegicus Procollagen II alpha 1 (Col2a1), mRNA	RC0-ST0379-210100-032-g04 ST0379 Homo sapiens cDNA	Interleukin-12 p35 subunit (mice, Genomic, 700 nt. segment 4 of 5)	HSC1RF022 normalized Infant brain cDNA Homo sapiens cDNA clone c-1rf02 3'	Mus musculus calcium channel, voltage-dependent, T type, alpha 1G subunit (Cacna1g), mRNA	601308679F1 NIH_MGC_44 Homo sepiens cDNA clone IMAGE:3827086 5	C.reinhardtil nuclear gene on linkage group XIX	A.immersus gene for transposase	G.gallus gene encoding non-histone chromosomal protein HMG-14b, exons 4 and 5	MR3-ST0290-280100-025-g07 ST0290 Homo saplens cDNA	MR3-ST0290-290100-025-g07 ST0290 Homo sapiens cDNA
Top Hit Database Source	L	EST_HUMAN	ΙN	SWISSPROT	N _T	Ę	EST_HUMAN	EST_HUMAN	ΝΤ	SWISSPROT	EST_HUMAN	EST_HUMAN		П	T_HUMAN	NT	EST_HUMAN	NT	EST_HUMAN	NT	NT	EST HUMAN	NT	EST_HUMAN	TN	EST_HUMAN	NT	NT	NT		EST_HUMAN
Top Hit Acession No.	M65109.1	1.2E-01 AV658033.1	1.2E-01 AJ271736.1	Q04912	AF039442.1		BE061418.1	A1299903.1	L10187.1	096433	1.2E-01 BF314481.1	AI561003.1		AA569006.1	BF697308.1	AL161560.2	AW972158.1	D64004.1	AU140363.1	6755215 NT	8978678 NT	AW821909.1	S82418.1	F03265.1	6753231 NT	BE393186.1	X62135.1	Y07695.1	X52708.1	AW819412.1	AW819412.1
Most Similar (Top) Hit BLAST E Value	1.2E-01	1.2E-01	1.2E-01	1.2E-01		1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.1E-01			1.1E-01	1.1E-01	1.1E-01	1.1E-01	1.1E-01	1.1E-01	1.1E-01	1.1E-01	1.1E-01	1.1E-01	1.1E-01	1.1E-01	1.1E-01	1.1E-01	1.1E-01	1.1E-01	1.1E-01
Expression Signal	2.02	222	2.78	2.58	7.69	1.41	2.36	3.68	2.07	4.72	2.18	0.95		3.38	1.53	1.29	4.08	1.72	2.47	2.25	1.17	1.17	1.84	8.0	1.39	3	1.54	0.94	1.23	1.31	1.31
ORF SEQ ID NO:				24897				25231				20301			20789		20839	20980	21267					22717		23082	23123	23246	23348		23699
Exon SEQ ID NO:	18612	18822	19043	19693	!	19230	19706	19281	19295	19836	19424	10483	<u> </u>			10985	12688	11137	11408	12150	12663	12394		L	13221	13293	13322	13460	13562	13925	13925
Probe SEQ ID NO:	8788	9032	9383	9458	9567	9671	9739	9761	9782	9786	8880	295		983	1038	1069	1143	1229	1504	2266	2492	2520	2825	2897	3280	3374	3405	3534	3648	4021	4021

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ORF SEQ ID NO:	Ø	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
13930		0.87	1.1E-01	AF030001.1	FN	Mus muscultus major histocompatibility locus class III region.butyrophilin-like protein gene, partial cds; Notch4, PBX2, RAGE, lysophatidic acid acyt transferase-alpha, palmitoyl-protein thioesterase 2 (PPT2), CREB-RP, and tenascin X (TNX) genes, compley
14069		7.83	1.1E-01	AF157068.1	Z	Drosophila melanogaster klarsicht protein (klar) mRNA, complete cds
14089 2	23867		1.1E-01	AW802056.1	EST_HUMAN	IL5-UM0070-020500-068-a08 UM0070 Homo sapiens cDNA
14360 2	24150	0.81	1.1E-01	AF064564.2	ΙΝ	Fugu rubripes neurofibrometosis type 1 (NF1), A-kinase anchor protein (AKAP84), BAW protein (BAW), and WSB1 protein (WSB1) genes, complete cds
	24214	2.02	1.1E-01	S44957.1	Į.	Tapa-1=integral membrane protein TAPA-1 [mice, B cell lymphoma line 38C13, Genomic, 1973 nt, segment 1 of 7]
	24397		1.1E-01		TN	A.immersus gene for transposase
14702	24487		1.1E-01		NT.	Synechocystis sp. PCC6803 complete genome, 10/27, 1188886-1311234
15399		1.43	1.1E-01	AA747216.1	EST_HUMAN	nx76a03.s1 NCI_CGAP_Ew1 Homo saplens cDNA clone IMAGE:1268140 similar to contains Alu repetitive element.contains element MER35 repetitive element.
15478	25551	1.54	1.1E-01	X68851.1	N T	S.pombe ste8 gene encoding protein kinase
15494	25570	4.73	1.1E-01	M86533.1	Į.	Providencia rettgari penicillin G amidase gene
15581	25681	1.46	1.1E-01	AJ007973.1	Z.	Homo saplens LGMD2B gene
16596	25697	1.79	1.1E-01	BE769162.1	EST_HUMAN	PM3-FT0024-130600-004-f12 FT0024 Homo saplens cDNA
15606	25708	7.01	1.1E-01	AW853699.1	EST_HUMAN	RC3-CT0254-280999-011-a01 CT0254 Homo sapiens cDNA
15782	25903	1.39	1.15-01	AF035746.1	EST_HUMAN	AF035746 Human salivary gland cell line HSG Homo sapiens cDNA clone RL43
15890	26012	3.48	1.1E-01	069635	SWISSPROT	ACETYL-COENZYME A SYNTHETASE (ACETATE-COA LIGASE) (ACYL-ACTIVATING ENZYME)
		2.9		AF032922.1	NT	Homo sapiens syntaxin 4 binding protein UNC-18c (UNC-18c) mRNA, complete cds
16052	26198	2.21	1.1E-01	11432372 NT	INT	Homo saplens phosphattdylinositol glycan, class B (PIGB), mRNA
16309	26474	7.05	1.1E-01	BF684628.1	EST_HUMAN	602140976F1 NIH_MGC_46 Homo sapiens cDNA clone IMAGE:4302019 5'
16309	26475	7.05	1.1E-01	BF684628.1	EST_HUMAN	602/140978F1 NIH_MGC_46 Homo sepiens cDNA clone IMAGE:4302019 5'
16372	26550	1.74	1.1E-01	P41067	SWISSPROT	TRAB PROTEIN
18300	26670	307	70 A 7	1 76266ZVV	EST LIMAN	ah31b08.s1 Soares_parathyroid_tumor_NbHPA Homo sapiens cDNA clone 1240403 3' similar to gb:J03483 CHROMOGRANIN A PRECLIRSOR /HIMAN)
16858	26845		1.1E-01		EST HUMAN	nh04g10.s1 NCI CGAP Thy1 Homo sapiens cDNA clone IMAGE:943362
16658	28846		1.15-01		EST HUMAN	nh04g10.s1 NCI_CGAP_Thy1 Homo saplens cDNA clone IMAGE:943362
16682	26872	1.22	1.1E-01	X91233.1	N	H.sapiens IL15 gene
16705		1.24	1.1E-01	AW817918.1	EST_HUMAN	PM1-ST0270-080200-001-f09 ST0270 Homo sepiens cDNA
16741	26934	1.78	1.1E-01	AL134349.1	EST_HUMAN	DKFZp547P194_r1 547 (synonym: htbr1) Homo saplens cDNA clone DKFZp547P194 5'
16995	27186	2.19	1 1E-01	U02482.1	Ę	Pediococcus acidiactici H plasmid pSMB74 pediocin AcH production (pap) gene cluster papA, papB, papC and papD genes, complete cds
l			l			

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Table 4
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			Г		T	т-				_		т-	_			Γ	_		1	Т	ī	\neg		_		_				ГТ	
	Top Hit Descriptor	zp93b12.r1 Stratagene muscle 937209 Homo sapiens cDNA clone IMAGE:627743 5	zp93b12,r1 Stratagene muscle 937209 Homo sapiens cDNA clone IMAGE:627743 5	yd19h03.s1 Soares fetal liver splean 1NFLS Homb sapiens cDNA clone IMAGE:108725 3' similar to gb://db.//db.//db.//db.//db.//db.//db.//d	MR2-GN0027-040900-005-e08 GN0027 Homo sapiens cDNA	y98a09.s1 Soares placenta Nb2HP Homo sapien; cDNA clone IMAGE:147064 3'	HSC1RF022 normalized infant brain cDNA Homo saplens cDNA clone c-1rf02 3'	Carassius auratus activin beta A precursor, mRNA, complete cds	yh35112.r1 Soares placenta Nb2HP Homo saplens cDNA clone IMAGE:131759 5' similar to contains Alu repatitive element.contains TAR1 repetitive element;	M.musculus cytokine gene	Z.mobilis tgt and lig genes encoding IRNA guanine transglycosylase and DNA ligase	Z.mobilis tgt and lig genes encoding tRNA guanine transglycosylase and DNA ligase	SKIN SECRETORY PROTEIN XP2 PRECURSOR (APEG PROTEIN)	RC2-NT0112-120600-014-f03 NT0112 Homo sepiens cDNA	601680551R2 NIH_MGC_83 Homo sapiens cDNA clone IMAGE:3950604 3'	601906350F1 NIH_MGC_54 Horno sapiens cDNA clone IMAGE:4134085 51	DEOXYRIBONUCLEASE II PRECURSOR (DNASE II) (ACID DNASE) (LYSOSOMAL DNASE II)	ws08d01.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2496577 3' similar to contains MER7.t3 MER7 repetitive element:	Arabidopsis thallana DNA chromosome 4, contig fragment No. 16	UI-H-BI3-alc-d-07-0-UI.s1 NCI_CGAP_Sub5 Homo saplens cDNA clone IMAGE:2736420 3'	601456301F1 NIH_MGC_66 Homo sapiens cDNA clone IMAGE:3859849 5	601906489F1 NIH_MGC_54 Homo sapiens cDNA clane IMAGE:4134071 5	QV2-NT0048-160800-316-e05 NT0048 Homo saplens cDNA	Chlamydophila pneumonlae AR39, section 91 of 94 of the complete genome	an32c04.y5 Gessler Wilms tumor Homo sapiens cDNA clone IMAGE:1700358 5'	Drosophila melanogaster tyrosine kinase p45 isoform (fer) mRNA, complete cds	EST364414 MAGE resequences, MAGB Homo sepiens cDNA	Mouse FTZ-F1 gene	UI-H-BW1-aca-e-12-0-UI.s1 NCI_CGAP_Sub7 Hcimo sapiens cDNA clone IMAGE:3084023 3'	zh62h04,s1 Soares_fetal_liver_spleen_1NFLS_S1/Homo sapiens cDNA clone IMAGE:416695 3'	Homo sapiens growth factor receptor-bound protein 7 (GRB7) gene, complete ods
		EST_HUMAN		EST HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	NT	EST HUMAN	PA	N	LN	SWISSPROT	EST_HUMAN	EST_HUMAN	EST_HUMAN	SWISSPROT	EST HUMAN	L L	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	TN	EST_HUMAN	TN	EST_HUMAN	TN	EST_HUMAN	EST_HUMAN	NT
	Top Hit Acession No.	AA192153.1	AA192153.1	172675.1		R80590.1	F03265.1	AF169032.1	R23708.1	X70058.1	Z11910.1	Z11910.1	P17437	BE767023.1	BE974556.1	BF239753.1	062855	A1985499.1	AL161504.2	AW451365.1	BF033991.1	BF239818.1	BF365703.1	AE002265.2	AI792349.1	U50450.1	AW952344.1	D49683.1	BF515935.1		AF274875.1
Most Similar	(Top) Hit BLAST E Value	1.15-01	1.1E-01	1.1E-01	1.16.01	1.15-01	1.1E-01	1.15-01	1.1E-01	1.1E-01	1.1E-01	1.15-01	1.1E-01	1.1E-01	1.15-01	1.15-01	1.0E-01	1.0F-01	1.0E-01	1.0E-01	1.0E-01	1.0E-01	1.0E-01	1.0E-01	1.0E-01	1.0E-01	1.0E-01	1.0E-01	1.0E-01	1.0E-01	1.0E-01
	Expression Signal		2.24	2.48	. 2.08	1.23	48.	3.88	2.83	1.95	3.21	3.21	2.79	3.19	2.08	2.14	4.08	1.89	2.88	-	96.0	1.01	2.44	1.5	1.17	1.26	2.06	0.87	1.44	9.18	11.21
	ORF SEQ ID NO:		27295	27362			22717		28530	28655	28676	28677	28763			25192		24007			23189	23359	23574	23987		24289	24502		24815		25680
	Exan SEQ ID NO:	17105	17105	17163	L	17711	12925	18173	18278	18391	18411	Į.	18491	18956	19507	19372	11094	11158	L	L	13384	13572	13782	14204	14349	14501	ł	15025	15051		15580
	Probe SEQ ID NO:	7228	7228	7287	7487	7861	8181	8294	8402	8519	8539	8539	8626	9241	9485	9893	1183	1261	1370	2439	3468	3658	3871	4307	4456	4613	4838	5158	5188	5260	5670

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Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acessian No.	Top Hit Datebase Source	Top Hit Descriptor
4131	14031	23805	6.67	9.8E-02	AF257329.1	N	Leptosphaeria maculans beta-tubulin mRNA, corriptete cds
4131			6.67	9.8E-02	AF257329.1	NT	Leptosphaeria maculans beta-tubulin mRNA, complete cds
8755	17904		2.1	9.8E-02	BF037421.1	EST HUMAN	601460793F1 NIH_MGC_66 Homo sepiens cDNA clone IMAGE:3864287 5'
1328	11235		1.16	9.7E-02	AB005808.1	NT	Aloe arborescens mRNA for NADP-malic enzyme, complete cds
1567	11471		0.98	9.7E-02	4503710 NT	ΙΝ	Homo seplens fibroblast growth factor receptor 3 (achondroplasia, thanatophoric dwarfism) (FGER3) mRNA
2214		22004	2.37	9.7E-02	BE168660.1	EST_HUMAN	QV1-HT0516-070300-095-e04 HT0516 Homo septens cDNA
3901			3.56	9.7E-02	Q99795	SWISSPROT	CELL SURFACE A33 ANTIGEN PRECURSOR (GLYCOPROTEIN A33)
5661	15572	25869	1.39	9.7E-02	AW954476.1	EST_HUMAN	EST366546 MAGE resequences, MAGC Homo sapiens cDNA
6321	16184		4.27	9.7E-02	Z99119.1	Ŋ	Bacillus subtilis complete genome (section 16 of 21): from 2997771 to 3213410
6855	16535		1.59			EST_HUMAN	yw41c03.s1 Weizmann Olfactory Epithelium Horrio sapiens cDNA clone IMAGE.254788 3'
6655	16535	26732	1.59	9.7E-02	N22798.1	EST_HUMAN	yw41c03.s1 Weizmann Olfactory Epithelium Homio sapiens cDNA clone IMAGE.254788 3'
7137	17014	27207	132	20-37 P	A1953984 1	FST HIMAN	wx78b08.x1 NCI_CGAP_Ov38 Homo sapiens cDNA clone IMAGE:2549747 3' similar to gb:X52851_ma1 PEPTIDYI -PROLYT CIS.TRANS ISOMERASE A /HI IMAN):
8531	18403		1.97			N	Mus musculus ligatin (Lgtn) mRNA, partial cds
1969	11862	21763	1.27	9.6E-02	Al080721.1	EST HUMAN	oz47d11.x1 Soares NhHMPu_S1 Homo sapiens cDNA clone IMAGE:16784853'
1969	11862	21764	127	9.6E-02	9.6E-02 AI080721.1	EST_HUMAN	oz47d11.x1 Soares_NhHMPu_S1 Homo sapiens conA clone IMAGE:1678485 3'
4248	14147	23921	6.02	9.6E-02	Z32686.2	N.	Proteus mirabilis fimbrial operon, strain HI4320
4932	14810	24579	0.94	0.6E-02	AW866230.1	EST HUMAN	EST378303 MAGE resequences, MAGI Homo sapiens cDNA
5093	14963	24738	0.8	9.6E-02	BE081729.1	EST_HUMAN	RC5-BT0254-031099-011-a03 BT0254 Homo sapiens cDNA
5713	15621		2.72	9.6E-02	BE910039.1	EST_HUMAN	601498088F1 NIH_MGC_70 Homo sapiens cDNA clone IMAGE:3900165 5'
7502	17371	27580	1.51		9.6E-02 AV687898.1	EST_HUMAN	AV687898 GKC Homo saplens cDNA clone GKCAAH02 5'
7677	17527		1.84	9.6E-02	BE894895.1	EST_HUMAN	601434080F1 NIH_MGC_72 Homo sapiens cDNA clone IMAGE:3919363 5'
7772	17622	27855	1.75		9.6E-02 AJ243211.1	TN	Homo sapiens DMBT1 candidate tumour suppressor gene, exons 1 to 55
7772	17622	27856	1.75		9.6E-02 AJ243211.1	L	Homo capiens DMBT1 candidate tumour suppressor gene, exons 1 to 55
7839	17689	27933	1.59		9.6E-02 AB013985.1	L	Antirrhinum majus transposon Tam3 pseudogene (for transposase (in S-5 copy)
7839	17689	27834	1.59	9.6E-02	9.6E-02 AB013985.1	INT	Antirrhinum majus transposon Tam3 pseudogene for transposase (in S-5 copy)
7886	17736	27980	3.29	9.6E-02		SWISSPROT	COMPLEMENT DECAY-ACCELERATING FACTOR PRECURSOR (CD55)
8125	18013	28260	6.51	9.6E-02	279702.1	NT	Mycobacterium tuberculosis H37Rv complete genome; segment 102/162
8957	18764	29056	1.8	9.6E-02	9.6E-02 AA625755.1	EST_HUMAN	zu91g01.s1 Soares_testis_NHT Homo saplens cDNA clone IMAGE:745392 3'
9798	19312		1.38	9.6E-02	9.6E-02 H14599.1	EST_HUMAN	ym19h03.s1 Soares infant brain 1NIB Homo capiens cDNA clone IMAGE:48653 3'
9848	19344	1	1.28		BE728219.1	EST HUMAN	601563355F1 NIH_MGC_20 Homo sapiens cDNA clone IWAGE:3832908 5'
4012	13918		2.25		9.5E-02 AW992395.1	EST_HUMAN	CM2-BN0023-050200-087-f12 BN0023 Homo sapiens cDNA
2062	14932	24704	0.87		U63374.1	N-	Lycopersicon esculentum polygalacturonase isoenzyme 1 beta subunit gene, complete cds

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Top: Hit Descriptor	Trimeresurus flavoviridis DNA for phospholipase A2 inhibitor, complete cds	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 38	601453642F1 NIH_MGC_66 Homo saplens cDNA clone IMAGE:3857243 5'	601453642F1 NIH_MGC_66 Homo sepiens cDNA clone IMAGE:3857243 5'	601453642F1 NIH_MGC_68 Homo sapiens cDNA clone IMAGE:3857243 5'	601453642F1 NIH_MGC_66 Homo saplens cDNA clone IMAGE:3857243 5'	602150882F1 NIH_MGC_81 Homo sapiens cDNA clone IMAGE:4291917 6:	Cavia porcellus 3beta-hydroxysteroid sulfotransferase mRNA, complete cds	Cavia porcellus 3beta-hydroxysteroid sulfotransferase mRNA, complete cds	M.capricolum DNA for CONTIG MC073	Mus musculus coding region determinant-binding protein (Crdbp), mRNA	Acinetobacter sp. cysD, cobQ, sodM, lysS, rubA, rubB, estB, oxyR, ppk, mtgA, ORF2 and ORF3 genes	Human BRCA1, Rho7 and vati genes, complete cds, and ipf35 gene, partial cds	Rattus norvegicus calcium channel alpha-1C subunit (ROB2) mRNA, partial cds	Human pephBGT-1 betaine-GABA transporter mRNA, complete cds	Homo sapiens BAI1-associated protein 3 (BAIAP3) mRNA	Homo sapiens nasopharyngeal epithelium specific protein 1 (NESG1), mRNA	602133086F1 NIH_MGC_81 Homo sapiens cDNA done IMAGE:4288269 5'	601286082F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3607653 5	601286082F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3607653 5'	AV732224 HTF Homo sapiens cDNA clone HTFAUA06 5	601655988R1 NIH_MGC_66 Homo saplens cDNA clone IMAGE:3855981 3'	HYPOTHETICAL PROTEIN KIAA0032	HYPOTHETICAL PROTEIN KIAA0032	Ul-H-811-afx-h-05-0-Ul.s1 NCI_CGAP_Sub3 Horrio sapiens cDNA clone IMAGE:2723553 3'	Photobacterium damselae subsp. damselae partial gyrB gene for DNA gyrase B subunit	hd28h12.x1 Soares_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:2910887 3'	Mus musculus major histocompatibility locus class II region; Fas-binding protein Dax (DAXX) gene, partial	cds; Bing1 (BING1), tapasin (tapasin), RalGDS-likė factor (RLF), KE2 (KE2), BING4 (BING4), beta1, 3-	galaciosy, transferase (beta 1,3-galactosy) tr>	Molluscum contagiosum virus subtype 1, complete genome	Molluscum contagiosum virus subtype 1, complete genome	Molluscum contagiosum virus subtype 1, complete genome
Top Hit Database Source	NT	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	N	Ŋ	NT	ΙN	LX.	LZ	L	NT	LN L	EST_HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	SWISSPROT	SWISSPROT	EST_HUMAN	LZ L	EST_HUMAN		!	LN L	L	Į.	NT
Top Hit Acession No.	AB003473.1	AL181538.2	BF035861.1	BF035861.1	BF035861.1	BF035861.1	BF671063.1	U55944.1	U56944.1	Z33059.1	0753517 NT	Z46863.1	L78833.1	U31815.1	U27699.1	4809280 NT	6912525 NT	BF575511.1	BE391943.1	BE391943.1	AV732224.1	BE962631.2	Q15034	Q15034	AW206117.1	AJ249850.1	AW468850.1			-			U60315.1
Most Similar (Top) Hit BLAST E Value			9.5E-02				9.4E-02	9.4E-02	9.4E-02	9.4E-02	9.4E-02	9.4E-02	9.4E-02	9.4E-02	9.4E-02	9.3E-02	9.3E-02	9.3E-02	9.3E-02		9.3E-02	9.3E-02	9.3E-02	9.3E-02 (9.3E-02	9.3E-02	9.3E-02/			9.3E-02/	9.2E-02 (9.2E-02	9.2E-02
Expression Signal	3.72	7.48	2.59	2.69	3.29	3.29	3.86	0.86	0.86	4.34	0.89	2.62	2.69	3.36	1.42	1.83	5.39	2.03	3.51	3.51	1.88	2.24	3.52	3.52	3.74	1,85	8.62			1.95	4.32	4.32	4.32
ORF SEQ ID NO:		28494				28207	21565		21598		24621		26505		25182				23738		_	27665	27948	27950					•		20008	20008	20010
Exon SEQ ID NO:	16186	16327				17957	11690	11717	11717	13716	14855	16876	16338	19603	19411	12887	12921	13142	13962	13962	14516	17450	17705	17705	17763	19527	19555			- 1	10197	10197	10197
Probe SEQ ID NO:	6323	6467	6233	6293	8088	8068	1792	1820	1820	3804	4980	6869	8304	9083	8843	2960	2993	3218	4060	4060	4628	7589	7855	7855	7913	9343	9704			888	87	228	228

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					7::-		
Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No	Top Hit Database Source	Tóp Hit Descriptor
2180	12087		22	9.2E-02	R54156.1	EST_HUMAN	yg98f07.r1 Soares infant brain 1NIB Homo sapiens cDNA clone IMAGE:41618 5'
3142	13067	L	65.4	9.2E-02	Q28631	SWISSPROT	MAJOR EPIDIDYMIS-SPECIFIC PROTEIN E4 (EPIDIDYMAL PROTEIN BE-20)
3265	13188	22986	1.16	9.2E-02	AA534354.1	EST_HUMAN	nf79e01.s1 NCI_CGAP_Co3 Homo sepiens cDNA clone IMAGE:926136 3'
3537	13453		1.12	9.2E-02	6755215 NT	님	Mus musculus pre T-cell antigen receptor alpha (Ptora), mRNA
4145	14045		1.34	9.2E-02	U92048.1	F	Human herpesvirus 1 strain KOS-63, latency-associated transcript, promoter region
4211	14109		1.02	9.2E-02	BE299722.1	EST_HUMAN	600944365F1 NIH_MGC_17 Homo saplens cDNA clone IMAGE:2960176 5
4538	14431	24213	1.86	9.2E-02	X96402.1	IN	G.gallus Mia-CK gene
6670	16550	26746	. 1.86	9.2E-02	T49920.1	EST_HUMAN	ye99c09.r1 Stratagene placenta (#937225) Homb sapiens cDNA clone IMAGE:69808 5' similar to similar to gb.X58009 GUANINE NUCLEOTIDE-BINDING PROTEIN G(S), ALPHA SUBUNIT (HUMAN)
6756	16635	26823		9.2E-02	X95256.1	N.	H. vulgare xylose Isomerase gene
417	9984		2.77	9.1E-02	X77665.1	TN	O. cuniculus k12 keratin gene
2365		22139	1.01	9.1E-02	P78985	SWISSPROT	6-PHOSPHOFRUCTOKINASE (PHOSPHOFRUCTOKINASE)
3618	13532		1.14	9.1E-02	AW372569.1	EST_HUMAN	PM2-BT0349-161299-001-f02 BT0349 Homo saplens cDNA
4383	14279	24058	1.81	9.1E-02	AL161554.2	NT	Arabidopsis thaliana DNA chromosome 4, contigifragment No. 64
2099	15426	26487	1.73	9.1E-02	AF129768.1	<u> </u>	Homo sapiens MSH55 gene, partial cds; and CLIC1, DDAH, G6b, G6c, G5b, G6d, G6e, G6f, BAT5, G5b, CSK2B, BAT4, G4, App M, BAT3, BAT2, AIF-1, 107, LST-1, LTB, TNF, and LTA genes, complete cds
6372	L	Ĺ	11.89		AW160658.1	EST_HUMAN	au74a05.y1 Schneider fetal brain 00004 Homo sapiens cDNA clone IMAGE:2781968 5'
7958			1.65	9.1E-02	T02984.1	EST HUMAN	FB19F10 Fetal brain, Stratagene Homo sapiens cDNA clone FB19F10 3'end
9021	18815		1,29	9.1E-02	9633494 NT	1. '	Bacteriophage Mu, complete genome
9226	19692		1.52	9.1E-02	AA179901.1	EST_HUMAN	മാ8h12.s1 Stratagene muscle 937209 Homo sapiens cDNA clone IMAGE:611783 3' similar to SW:TRT3_HUMAN P49378 TROPONIN T, FAST SKELETAL MUSCLE, ISOFORM BETA ;
9785	19548		6.63	9.1E-02	AJ291390.1	NT	Homo saplens partial MUC3B gene for MUC3B mucin, exons 1-11
			,				FOLATE RECEPTOR ALPHA PRECURSOR (FR-ALPHA) (FOLATE RECEPTOR 1) (FOLATE RECEPTOR, ADULT) (ADULT FOLATE-BINDING PROTEIN) (FBP) (OVARIAN TUMOR-ASSOCIATED
727	10659	20490	3.36	9.0E-02	P15328	SWISSPROT	ANTIGEN MOV18) (KB CELLS FBP)
4044			27 0	00 00	4 60700020	MANIET TOT	hV39g10.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3175842 3' similar to contains Alu
	_			8.05-02		ביים ביים	Table and the second of the se
2772	_!			9.0E-02	AF138522.1	Z	HIV-1 pscuss-us from USA envelope glycoprotein (env) gene, partial cas
2772			1.41	9.0E-02	AF138522.1	LN	HIV-1 p8c095-06 from USA envelope glycoprotein (env) gene, partial cds
3294	1		0.92		AF279135.1	Ę	Dictyostelium discoldeum spore coat structural protein SP65 (cotE) gene, complete cds
4202	- 1					N	corticosteroid-binding globulin [Saimiri sciureus≕squirrel monkeys, liver, mRNA, 1474 nt]
4202	_1			9.0E-02	S68757.1	뉟	corticosteroid-binding globulin [Saimiri sciureus≂stjuirrel monkays, liver, mRNA, 1474 ntj
4321	14218	24001	1.2	9.0E-02	P55268	SWISSPROT	LAMININ BETA-2 CHAIN PRECURSOR (S-LAMININ)

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	Top Hit Descriptor Source	NT Plasmodium falciparum P-type ATPase 3 gene	SWISSPROT REGULATORY PROTEIN ZESTE	za88a12.r1 Soares_fetal_lung_NbHL19W Homo sapiens cDNA clone IMAGE:297694 5' similar to lect. In IMAN PIR-SE2171 SE2171 small G protein - human	Т	EST HUMAN	EST HUMAN		.1 NT Atrichum engustatum AtranFlo2 protein (AtranFlò2) gene, partial cds	EST_HUMAN	EST_HUMAN	Γ	11433478 NT Homo sapiens similar to endoglycen (H. sapiens) (LOC63107), mRNA	FOLD BIFUNCTIONAL PROTEIN [INCLUDES:METHYLENETETRAHYDROFOLATE SWISSPROT DEHYDROGENASE; METHENYLTETRAHYDROFOLATE CYCLOHYDROLASE]		.1 EST_HUMAN EST180187 Liver, hepatocellular carcinoma Homo capiens cDNA 6' end	.1 EST_HUMAN 902129882F1 NIH_MGC_56 Homo sepiens cDNA clone IMAGE:4286180 5	SWISSPROT PROBABLE DNA LIGASE (POLYDEOXYRIBONUCLEOTIDE SYNTHASE (ATP))	EST_HUMAN	TRANSCRIPTION INITIATION FACTOR TFIID 135 KDA SUBUNIT (TAFIL 135) (TAFIL 135) (TAFIL 130)		EST_HUMAN	.1 EST_HUMAN 601191770F1 NIH_MGC_7 Homo septens cDNA clone IMAGE:3535648 5'	.1 EST_HUMAN 601191770F1 NIH_MGC_7 Homo septiens cDNA clone IMAGE:3535648 5'	1 EST_HUMAN DKFZp434D1313_r1 434 (synonym: hies3) Home sapiens cDNA clone DKFZp434D1313 5	NT S.cerevislae chromosome XIV reading frame ORF YNL285w	Homo sapiens zinc finger protein 92 (ZFP92), expressed-Xq28STS protein (XQ28ORF), and biglycan (BGN) genes, complete cds; and plasma membrane calcium ATPase isoform 3 (PMCA3) gene, partial cds	
	#F 99	Plasmodium falciparum P-type A			T	1	Г		Atrichum angustatum AtranFlo2 p			Г	Homo sapiens similar to endoglyc			П	_		Γ	Π.	Т	П	Γ	Г		П	Homo sapiens zinc finger protein genes, complete cds, and plasma	(IAOG) en allall ben (3GOGCA), sistem STOGCA benonium (COGOZ) CO sistem comit enim ancient amol
,		뉟	SWISSPR	1 I	IN 65	EST HUM	EST HUM	EST_HUM	TN	EST_HUM	EST_HUM	EST_HUM	78 NT	SWISSPR	ž	EST_HUM	EST_HUM	SWISSPR	EST_HUM	GIAICCOD	NT	EST_HUM	EST_HUM	EST_HUM	EST HUN	N	TN	
	Top Hit Acession No.	X65740.2	Q24597	W 56037 4	114317	BF7015	BF701593.1	BE153572.1	AF286055.1	AA424887.1	AW452122.1	AW452122.1		P47259	279021.1	AA309319.1	BF696918.1	027474	AA299128.1	980000		AA15187	BE264455.1	BE284455.1	AL040129.1	271561.1	U82695.2	
	Most Similar (Top) Hit BLAST E Value	9.0E-02	9.0E-02	0 0 50 0	9.0E-02	8.9E-02	8.9E-02	8.9E-02	8.9E-02	8.9E-02	8.9E-02	8.9E-02	8.9E-02	8.9E-02	8.9E-02	8.9E-02	8.9E-02	8.8E-02	8.8E-02	0 00 00	8.8E-02	8.8E-02	8.8E-02	8.8E-02	8.8E-02	8.8E-02	8.7E-02	
	Expression Signal	1.79	1.08	α	15.35	2.15	2.15	1.41	1.71	0.92	3.3	3.3	3.13	.58	1.83	5.28	4.03	1.25	96.0	70 6	1.13	1.8	3.19	3.19	10.63	1.38	3.02	
	ORF SEQ ID NO:	24248	24780	25553		21189				24208		L	25568	26290		27088		21113	23532			27280			28722		23337	
	Exon SEQ ID NO:	14460	15009	18580		L	ı	ł _	14004	14427	15485	15485	15491	16135	L	16875	18949	11257	13740	42050	1	17072	18334	18334	18453		13550	
1	Probe SEQ ID NO:	4568	6142	5647	9987	1418	1419	2338	4104	4534	5569	5569	5576	6270	6460	8669	9228	1351	3828	9706	4205	7195	8461	8461	8585	9302	3636	

· Page 89 of 413 Table 4 Single Exon Probes Expressed in Heart

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	Top Hit Descriptor	zh68a02.r1 Soares fetal liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:417194 5' similar to contains element MER12 repetitive element;	Mus musculus JNK interacting protein-3a (Jip3), mRNA, complete cds	Methanobacterium thermoautotrophicum from bases 1176181 to 1189406 (section 101 of 148) of the	complete genome	zs55g08.s1 NCI_CGAP_GCB1 Homo septens cDNA clone IMAGE:701438 3'	zs55g08.s1 NCI_CGAP_GCB1 Homo saplens cDNA done IMAGE:701438 3'	Oryctolagus cuniculus cytochrome P-450 (CYP4A4) gene, 5' end	Gluconobacter oxydans tRNA-lle and tRNA-Ala genes	Human DNA for immunoglobulin alpha heavy chain from a case of alpha heavy chain disease	Mus musculus nidogen 2 (Nid2), mRNA	Homo saplens Xq pseudoautosomal region; segrinent 2/2	601304016F1 NIH_MGC_21 Homo saplens cDNA clone IMAGE:3638643 57	Trichomonas vaginalis beta-tubulin (btub1) gene complete cds	Dictyostelium discoideum adenylyl cyclase (acrA) gene, complete cds	Oryctolagus cuniculus galectin-3 gene, untranslated exon and 5' flanking region	Homo capiens LCN1b gene	Mouse germline IgM chain gene, D region; D-q52, mu switch region (part a)	Mouse garmline IgM chain gene, D region; D-q52, mu switch region (part a)	Homo sapiens Snf2-related CBP activator protein (SRCAP) mRNA	Homo saplens Snf2-related CBP activator protein (SRCAP) mRNA	Lacerta media cytochrome c oxidase subunit 1 gene, partial cds; mitochondrial gene for mitochondrial product	Lacerta media cytochrome c oxidase subunit 1 gene, partial cds; mitochondrial gene for mitochondrial product	601893437F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:4139216 5	601893437F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:4139216 5	Archaeoglobus fulgidus section 34 of 172 of the complete genome	Bacillus stearothermophilus BsrFI methylase (FIM) and BsrFI restriction endonuclease (FIR) genes, complete	cds	Helicobacter pylori 28695 section 130 of 134 of the complete genome	M PROTEIN, SEROTYPE 6 PRECURSOR	Mus musculus phospholipase C-like protein mRNA, partial cds	Mus musculus myosin XV (Myo15), mRNA
	Top Hit Database Source	EST_HUMAN	NT		Ę	EST_HUMAN	EST_HUMAN	IN	IN	LN	TN	ΙN	EST_HUMAN	ĮŅ.	ΝΤ	TN	NT	L	IN	IN	TN	NT .	TN	EST HUMAN	EST HUMAN			L	μN	SWISSPROT	Ŋ	۲.
	Top Hit Acesslon No.	W87841.1	AF178636.1		AE000895.1	1	AA286875.1	L04758.1	AJ007763.1	X17116.1	6679057	AJ271736.1	BE408667.1	L05468.1	AF153362.1	U68179.1	Y10826.1	J00440.1	300440.1	5730066 NT	5730066 NT	AF206551.1	AF206551.1	BF305606.1	BF305606.1	AE001073.1		AF283660.1	AE000652.1	P08089	AF233885.1	6754779 NT
Mact Cimilor	(Top) Hit BLAST E	8.7E-02	8.7E-02				8.7E-02		8.7E-02	8.7E-02	8.7E-02	8.6E-02	8.6E-02	8.6E-02	8.6E-02	8.6E-02	8.6E-02	8.6E-02		8.6E-02	8.6E-02	8.6E-02	8.6E-02	8.6E-02		8.6E-02		8.6E-02	8.5E-02			8.5E-02
	Expression Signal	0.82	1.22		1.08	5.41	5.41	2.58	1.77	2.58	1.81	7.05	222	2.94	3.07	0.87	4.38	1.51	1.51	1.28	1.26	1.98	1.98	3.63	3.63	4.71		1.73	2.87	1.8	6.34	1.93
	ORF SEQ ID NO:	23578				24951	24962		28770			20991				24060	25718	25867	25868			28441	28442						22124			
	Exon SEQ ID NO:	13790					15177	17985	18496	18894	19105	11139	12084	13078	<u>1</u>	14281	15616		15752	16508	16508	18182	18192	L	1	ł	ļ	- 1	12227	1	1 1	16880
	Probe SEQ ID NO:	3878	4609		5034	5255	6265	8094	8631	9293	9484	1232	2197	3151	3593	4385	5708	5846	5846	6628	6828	8315	8315	8674	8574	8738		8888	2347	5500	9999	7003

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Single Exon Probes Expressed in Heart

Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Тор Hit Descriptor
7881	17511	27737	3.07	8.5E-02	BE833054.1	EST_HUMAN	RC4-OT0037-200700-014-e05 OT0037 Homo saplens cDNA
7661			3.07	8.5E-02	BE833054.1	EST_HUMAN	RC4-OT0037-200700-014-e05 OT0037 Homo sapiens cDNA
8498			11.15	8.5E-02		INT	Homo sapiens heparanase precursor, mRNA, complete cds
8512	18384	28649	4.07	8.5E-02	AB001562.1	NT	Streptococcus mutans gene for glucose-1-phosphate unidylytransferase, complete cds
9681	19488		1.39	8.5E-02	AJ005586.1	LZ LZ	Antirrhinum majus mRNA for MYB-related transcription factor
9841	19339		3.28	8.5E-02	AA362934.1	EST_HUMAN	EST72736 Ovary II Homo sapiens cDNA 5' end
2632	12728	22391	474	8.4E-02	W69330.1	EST_HUMAN	zd44e11.r1 Soares_fetal_heart_NbHH19W Homo sapiens cDNA clone IMAGE:343532 5'
5253					BE267153.1		601190436F1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:3534383 5'
6019	15923		1.7.1		AK024458.1	LN	Homo sapiens mRNA for FLJ00050 protein, partial cds
0899	16560	26755	8.11	8.4E-02	BE095074.1	EST_HUMAN	CM3-BT0790-260400-162-d05 BT0790 Homo sapiens cDNA
7934	17784	28020	1 44	8 4F-02	11735184 1	NAMI H TAR	as88910.x1 Barstead colon HPLRB7 Homo sapiens cDNA clone IMAGE:2335842 3' similar to TR:O88312 O88312 GOB-4
1085	1	L			Reagean	FN	Ivodes havanouis militorinondrin complete manoma
	L	1		0.3E-02			invocations introduction; complete general
1965	- 1				6836880	IN.	Ixodes nexagonus mitocnonditon, complete genome
3544	1				P75334	SWISSPROT	HYPOTHETICAL LIPOPROTEIN MG309 HOMOLOG PRECURSOR
3567	13481	17252	0.88		AI436797.1		th82g06.x1 Soares_NhHMPu_S1 Homo septens cDNA clone IMAGE:2125210 3'
3567	13481	23272	0.88		A1436797.1	EST_HUMAN	th82g06.x1 Soares_NhHMPu_S1 Homo capiens cDNA clone IMAGE:2125210 3'
5840	15746		2.82	8.3E-02	AF052683.1	IN	Homo saplens protocedherin 43 gene, exon 1
6653	16533	26728	3.42		AF185787.1	IN	Rattus norvegicus dystrophin-related protein 2 A-form splice variant (Drp2) mRNA, complete cds
	L						og88g08.s1 NOL_CGAP_Kid5 Homo sapiens cDNA clone IMAGE:1455422 3' similar to contains L1.t1 L1 L1
6871			1.47	8.3E-02	AA865285.1	EST_HUMAN	repetitive element;
6829	16708		1.42	8.3E-02	AA987873.1	EST_HUMAN	oq81f10.s1 NCI_CGAP_Kid6 Hamo sapiens cDNA clane IMAGE:1592779 3'
7408	88221	97579	77 }	CO-36 8	A1M 583503 1	NAMILE TOO	1805h10.x1 Human Pancreatic Islets Homo septems cDNA 3' similar to TR:Q16332 Q15332 GAMMA SUBLINIT OF SODILIM POTASSII IM A TPASS LIIKE
7508			18		Γ	LN	Arabidopsis thallana DNA chromosome 4, contig fragment No. 91
9307	L		1.55		Γ	EST HUMAN	601644770F1 NIH MGC_56 Homo sapiens cDNA clone IMAGE:3929993 5'
1357	11263		7.82		Y08170.2	L	Gallus gallus mRNA for for OBCAM protein gamma isoform
1480	11385	21248		8.2E-02	AF167077.2	ĽΝ	Canis familiaris glutamate transporter (EAAT4) mRNA, complete cds
3036	12964		1.78		AL.163206.2	NT	Homo saplens chromosome 21 segment HS21C006
3733	13645		1.26	8.2E-02	AL161498.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 10
3928	13835	23615	1.11	8.2E-02	AL163206.2	L	Homo sapiens chromosome 21 segment HS21C006
4187	14087		5.36		P48960	SWISSPROT	LEUCOCYTE ANTIGEN CD97 PRECURSOR
4187	14087			8.2E-02	P48960	SWISSPROT	LEUCOCYTE ANTIGEN CD97 PRECURSOR
4187	14087	23864	5.36	8.2E-02	P48960	SWISSPROT	LEUCOCYTE ANTIGEN CD97 PRECURSOR

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Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
8172	18060	28310	7.42	8.0E-02	AF217798.1	NT	Homo sapiens SCG10 like-protein, helicase-like protein NHL, M68, and ADP-ribosylation factor related protein 1 (ARFRP1) genes, complete ods
9344	19023	25300	2.94	8.0E-02	AJ005375.1	Į,	Drosophila orena hunchback region
9891	13887		1.47	8.0E-02	4503034	IN	Homo saplens cAMP responsive element binding protein-like 2 (CREBL2) mRNA
6966	19854		3.04	8.0E-02	AJ278435.1	TN	Mus musculus Ranbp7 gene, Staf gene and Wee1 gene
2127	12015	21913	3.98	7.9E-02	BE250008.1	EST_HUMAN	600943191F1 NIH_MGC_15 Hama saplens cDNA clane IMAGE:2959510 5
2048	12875	E7800	E7 9	20-36 Z	A 1582029 1	HAT HIMAN	ar98c08.x1 Barstead colon HPLRB7 Homo sepiens cDNA clone IMAGE:2173646 3' similar to gb:Z26876 605 RIBOSOMAL PROTEIN 138 (HUMAN):
3776							Mus musculus colony stimulating factor 1 receptor (Osf1r), mRNA
3776	<u> </u>		3.31	7.9E-02		N N	Mus musculus colony stimulating factor 1 receptor (Csf1r), mRNA
4595	14483		1.08	7.9E-02	BF34845	EST_HUMAN	602019770F1 NCI_CGAP_Brn67 Homo sapiens cDNA clone IMAGE:4155401 5'
4706	14592		1.16	7.9E-02	AB008019.1	N-	Arabidopsis thaliana RXW24L mRNA, partial cds
2899	16562	26756	3.25	7.9E-02	U27832.1	LN LN	Saccharomyces cerevisiae suppressor of MIF2 Smt4p (SMT4) gene, complete cds
	1 _					Г	ou63b05.s1 NCI_CGAP_Br2 Homo sapiens cDNA done IMAGE:1632465 3' similar to WP:C37A2.2
7762	17612	27838	5.68	7.9E-02	A1081644.1	EST_HUMAN	CE08811;
22.5	07027	COOLO		00 00 5		14711 11 120	ou63b05.s1 NCI_CGAP_Br2 Homo sapiens cDNA clone IMAGE:1632465 3' similar to WP:C37A2.2
1,02	1/0/1	1	0.00		ANO IOTH. I	NAWOL 181	1100000
1192	11102	20947	1.43	7.8E-02	AI793275.1	EST_HUMAN	occeduz. yo not journ't Lub momo sapiens cunna done image: 15/0407 o similiar to contains L1.13 L1 repetitive element ;
	[L			_		0059402.y6 NCI_CGAP_Lu5 Homo sepiens cDNA clone IMAGE:1570467 5' similar to contains L1.t3 L1
1192	_1				AI793275.1	EST_HUMAN	repetitive element;
4688		24371	0.81		BE836331.1	EST_HUMAN	PM3-FN0058-140700-005-f09 FN0058 Homo saplens cDNA
6019	13603		2.71	7.8E-02	BE250048.1	EST HUMAN	600943055F1 NIH_MGC_15 Homo sapiens cDNA clone IMAGE:2959693 5'
6197	15060	24825	1.04	7.8E-02	AI418520.1	EST_HUMAN	tg48g12.x1 Soares_NFL_T_GBC_S1 Homo saptions cDNA clone IMAGE:2112070 3' similar to contains MER10.t3 MER10 repetitive element ;
							Homo sapiens FWE domain-containing dual specificity protein phosphatase FWE-DSP1b mRNA, complete
7238	17113	27308	2.06	7.8E-02	AF233437.1	NT	spo
		,.					Homo sapiens FWE domain-containing dual specificity protein phosphatase FWE-DSP1b mRNA, complete
7236			2.06			NT	cds
6867	17307		1.27	7.8E-02	AA468354.1	EST_HUMAN	nc68b06.r1 NCI_CGAP_Pr1 Homo sepiens cDNA clone IMAGE:771731
1378	12693	21139	1	7.7E-02		NT	Homo sapiens WRN (WRN) gene, complete cds
3538	13454		2.09	7.7E-02	AJ238093.1	NT	Homo saplens partial AF-4 gene, exons 2 to 7 and Alu repeat elements
6615	16495	26682	4.4	7.7E-02	AA402949.1	EST_HUMAN	zu53d11.r1 Soares ovary tumor NbHOT Homo sapiens cDNA clone IMAGE:741717 5' similar to TR:G1173905 G1173905 SPLICEOSOME ASSOCIATED PROTEIN.;
				١			

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Probe SEQ ID NO: NO: 1660 8376 9835 3341 3341 1444 4407 7695 6855 6953 4605 4605 4605 6955 6955 6955 6955 6955 6955 6955 6	Exan SEQ ID NO: 17510 17510 13281 13281 13281 13281 13281 14650 14650 14650 14774 14301 16831 1439 14494 14573 14494 14573 14581 14581 14581 14581 16828	ORF SEQ ID NO: 27736 23081 23081 23081 23081 27024 24085 24085 24282 24282 24370 24370 24618 24618	Expression Signal Signal A 622 4 62 2 2 2 2 2 2 2 2 2 4 6 2 2 4 6 2 2 4 6 2 2 4 6 2 2 4 6 2 2 6 8 8 3 3 3 8 3 3 3 8 3 3 3 8 3	Most Similar (Top) Hit BLAST E Value Value 7.7E-02 7.7E-02 7.7E-02 7.6E-02 7.6E-02 7.6E-02 7.6E-02 7.6E-02 7.6E-02 7.6E-02 7.7E-02 p Hit Aces: No. No. No. 11422 11432 11432 11432 1143	Top Hit Database Source Source Source Source Source NT NT EST HUMAN EST HUMAN NT NT NT NT NT NT NT NT NT NT NT NT NT	Top Hit Descriptor Top Hit Descriptor Top Hit Descriptor PROBABLE SERNIETHREONINE-PROTEIN KINASE YBR059C Self-control of S	
9271	19675		2.08	7.4E-02	11525893 AW379431.1	NT EST_HUMAN	Homo sapiene histone deacetylase 6 (NY-CO-9), inRNA CM4-HT0243-081199-037-d11 HT0243 Homo sapiens cDNA
9678		25242	1.81	7.4E-02 7.3E-02	П	EST HUMAN	601453813F1 NIH_MGC_66 Homo saplens cDNA done IMAGE:3857738 5' 801658738R1 NIH_MGC_69 Homo saplens cDNA clone IMAGE:3886209 3'

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Single Exon Probes Expressed in Heart

-		_	_	_	_	_	_	_	_					-	_		_		_	_	_	_	_	_	_			
	Top Hit Descriptor	601658738R1 NIH_MGC_69 Homo sapiens cDNA clone IMAGE:3886209 3'	Themotoga maritima section 101 of 136 of the complete genome	CM0-NN1004-130300-284-g08 NN1004 Homo saplens cDNA	Homo sapiens chromosome 21 segment HS21¢102	Mus musculus transcription factor USF2 (USF2) gene, exons 8-10 and complete cds	PROLINE-RICH PROTEIN MP-3	PROLINE-RICH PROTEIN MP.3	Homo sepiens KIAA0424 protein (KIAA0424), mRNA	224802.s1 Soares_fetal_liver_spleen_1NFLS_\$1 Homo sapiens cDNA clone IMAGE:451178 3' similar to gb:L02426 26S PROTEASE SUBUNIT 4 (HUMAN);	Methanobacterium thermosutotrophicum from bases 1029155 to 1039934 (section 88 of 148) of the complete	garane	Methanobacterium thermoautotrophicum from bases 1029155 to 1039934 (section 88 of 148) of the complete genome	Homo sapiens chromosome 21 segment HS21C/101	Homo saplens chromosome 21 segment HS21C ₁ 101	Human Immunodeficiency virus type 1 isolate 26 reverse transcriptase (pol) gene, internal fragment, partial	UI-P-BWO-ail-a-05-0-1/1 s1 NCI CGAP Sub6 Home sablens cDNA clone IMAGE:2732049 3*	602077757F1 NIH MGC 62 Homo septens cDNA clone IMAGE:4251950 5'	Rhodomonas salina mitochondrion, complete genome	Streptococcus mutans gene for glucose-1-phosphate untdylytransferase, complete cds	Methanococcus jannaschii section 73 of 150 of the complete genome	CALMODULIN	601883558F1 NIH_MGC_57 Homo sapiens cDNA clone IMAGE:4095710 5'	Strongylocentrolus purpuratus mitochandrian, complete genome	AV712452 DCA Homo sapiens cDNA done DCAAUG01 5'	Homo saplens plasma membrane calcium ATPasia Isoform 1 (ATP2B1) gene, alternative splice products, partial cds	hq24f11.x1 NCI_CGAP_Adr1 Homo sapiens cDNA clone IMAGE:3120333 3' similar to TR:Q9Z340 Q9Z340 ATYPICAL PKC SPECIFIC BINDING PROTEIN.;	Homo sapiens zinc finger protein 92 (ZFP92), expressed-Xq28STS protein (XQ28ORF), and biglycan (BGN) genes, complete cds; and plasma membrane calcium ATPese isoform 3 (PMCA3) gene, partial cds
2	Top Hit Datebase Source	EST_HUMAN	L	EST_HUMAN	LN	N-	SWISSPROT	SWISSPROT	ΤN	EST_HUMAN		ż	Ę	Z	LZ	Ę	FST HIMAN	EST HUMAN	N	N	칟	SWISSPROT	EST_HUMAN	TN	EST_HUMAN	Į.	EST_HUMAN	Ę
•	Top Hit Acesslan No.	BE964961.2	AE001789.1	AW800281.1	AL163302.2	U12283.1	P05143	P05143	7662107 NT	AA779977.1		AE000882.1	AE000882.1	AL163301.2	AL163301.2	114704 4	AW 208322 1	BF572307.1	11466583 NT	AB001562.1	U67531.1	P11120	BF216086.1 EST_	5834897	AV712452.1	L14561.1	AW873187.1	U82695.2
	Most Similar (Top) Hit BLAST E Value	7.3E-02	7.3E-02	7.3E-02	7.3E-02				7.3E-02	7.3E-02		1.2E-02	7.2E-02	7.2E-02	7.2E-02	7 25.02	7 25.02	7.2E-02	7.2E-02		7.2E-02	7.2E-02		7.2E-02			7.2E-02	7.2E-02
	Expression Signal	96.0	2.73	3.04	14.81	1.01	2.44	2.44	1.27	2.78		0.84	0.94	2.23	2.23	c c	230	4.02	78.82	0.94	3.03	7.62	9.33	1.75	2.05	4.23	2.64	. 2.05
	ORF SEQ ID NO:	20223						26438		25910		1891	18912				23511	L	L		24922	24923			27608	27697	27818	28016
	Exon SEQ ID NO:			12695	12705	14806	16275	16275	16628	15788		10093	10093	L	11365	14661	Ł	L	L	L	15154	15155	16118		17396	17476	17596	17776
	Probe SEQ ID NO:	461	699	1485	1801	4927	6413	6413	6749	8548		-	114	1460	1460	2502	3840	4249	4594	4997	5230	5231	6252	6281	7646	7625	7746	7926

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Vatue	Top Hit Acession No.	Top Hit Database Source	Тър Hit Descriptor
7983	17833	28073	5.47	7.2E-02	BE565003.1	EST_HUMAN	601343928F1 NIH_MGC_63 Homo saplens cDNA clone IMAGE:3885951 5'
7894	17844		3.2	7.2E-02	BE539214.1	EST_HUMAN	601065194F1 NIH_MGC_10 Homo sapiens cDNA clone IMAGE:3451559 5'
8285	18164	28407	4.8	7.2E-02	AF049874.1	LN	Rattus norvegicus bHLH transcription factor Mist1 (Mist1) gene, complete cds
9178	18919	25346	1.44	7.2E-02	AA773696.1	EST_HUMAN	af81a04.r1 Soares_NhHMPu_S1 Homo sapiens cDNA clone IMAGE:1048398 5
9214	18942			7.2E-02		EST_HUMAN	AJ230796 Homo saplens library (Seranski P) Homo saplens cDNA clone PS13D5 3'
8273	18977		1.44	7.2E-02		EST_HUMAN	no05h08.s1 NCI_CGAP_Phe1 Homo sapiens cDNA clone IMAGE:1099839 3'
9332	19013		1.62	7.2€-02	U82828.1	NT	Homo sapiens ataxia telangiectasia (ATM) gene complete cds
9346	19531		3.89	7.2E-02	AW900962.1	EST_HUMAN	CM4-NN1009-200300-118-c11 NN1009 Homo sapiens cDNA
1862	11758	21633	1.65	7 15-02	1.02280.1	<u> </u>	Human immunodeficiency virus two 1 (D9) proviral structural cansid protein (gag) gene, partial cds
2243	L		0.9	7.1E-02		Į.	Pseudomonas aeruginosa PA01, section 451 of 529 of the complete genome
2247	_	22028	5.08	7.1E-02	-	EST HUMAN	601872281F1 NIH_MGC_53 Homo sapiens cDNA clone IMAGE:4092981 5
8063	18844		4.88	7.1E-02	BE304764.1	EST HUMAN	601143974F1 NIH_MGC_15 Homo saplens cDNA clone IMAGE:3051234 5'
517	10459	20270	1.23	7.0E-02	Q07092	SWISSPROT	COLLAGEN ALPHA 1(XVI) CHAIN PRECURSOR
1484	11389		1.46	7.0E-02	X96677.1	Ę	M.artiellia Mtcut-1 gene
1725	11626	21495	1.36	7.0E-02	AA056343.1	EST HUMAN	266f04.s1 Stratagene colon (#937204) Homo saplens cDNA clone IMAGE:5095993'
2994	12922	22715	1.72	7.0E-02	AW138152.1	EST_HUMAN	UI-H-BI1-acy-c-07-0-UI.s1 NCI_CGAP_Sub3 Homo sepiens cDNA clone IMAGE:2716020 3'
3822	13734	23523	-	7.0E-02	AA815438.1	EST_HUMAN	ai65a12.s.1 Soares_testis_NHT Homo sepiens cDNA clone 1375678.3' similar to gb:K03002 60S RIBOSOMAL PROTEIN L32 (HUMAN);
3958	13866	23842	1.24	7.0E-02	BE070264.1	EST HUMAN	QV4-BT0407-280100-090-e10 BT0407 Homo sapiens cDNA
4047	13949		1.08	7.0E-02	AW 792962.1	EST_HUMAN	CM0-UM0001-060300-270-e12 UM0001 Homo saplens cDNA
4121	14021	23799		7.0E-02	AF077821.1	TN	Canis familiaris Inducible nitric oxide synthase mRNA, complete cds
4846	14727	24510	1.7	7.0E-02	BF381987.1	EST_HUMAN	601816291F1 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:4050071 5'
7259	17136	82842	1.25	7.0E-02	9628113 NT	NT	African swine fever virus, complete genome
7534	17385	27697	1.19	7.0E-02	K02901.1	NT	Rat ig germline epsilon H-chain gene C-region, 3' end
8679	18567	28850	2.39	7.0E-02	AA724295.1	EST_HUMAN	ah99a05.s1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:1327184 3' similar to gb:L14837 TIGHT JUNCTION PROTEIN ZO-1 (HUMAN);
503	10445			6.9E-02	AL163210.2	L	Homo saplens chromosome 21 segment HS21C010
503		20257	5.34	6.8E-02	AL163210.2	NT	Homo capiens chromosome 21 segment HS21C010
1311	11217		1.31	6.9E-02	TN 8967057	TN	Homo sapiens regulator of Gz-selective protein signaling (ZGAP1) mRNA, and translated products
3724	13636	23421	1.42	6.9E-02	Q06364	SWISSPROT	26S PROTEASOME REGULATORY SUBUNIT S3 (NUCLEAR ANTIGEN 21D7)
3724	13636	23422	1.42	6.9E-02	Q06364	SWISSPROT	28S PROTEASOME REGULATORY SUBUNIT S3 (NUCLEAR ANTIGEN 21D7)
5100	14968		1.05	6.9E-02	AF079906.1	NT	Rables virus isolate b615 glycoprotein gene, partial¦cds

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Probe SEQ ID S NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acesslon No.	Top Hit Database Source	Top Hit Descriptor
6978	16855		1.37	6.9E-02	6.9E-02 BE567435.1	EST_HUMAN	601340661F1 NIH_MGC_53 Homo sapiens cDNA clone IMAGE:3883030 5'
8269	16855	27049	1.37	6.9E-02		EST_HUMAN	601340661F1 NIH_MGC_53 Homo sapiens cDNA clone IMAGE:3883030 5'
8208	18939		3.85	6.9E-02	6.9E-02 X74316.1	TN	X.laevis XFD2 mRNA for fork head protein
9385	19045		1.5	6.9E-02 P44821		SWISSPROT	PROTEIN TRANSPORT PROTEIN HOFC HOMOLOG
8658	19180		2.19	6.9E-02	6.9E-02 AF195953.1	INT	Homo sepiens membrans-bound aminopeptidase P (XNPEP2) gene, complete cds
1839	11738	21610	1.11	6.8E-02)	6.8E-02 AA496759.1	EST HUMAN	ae30f02.rl Gessler Wilms tumor Homo sapiens cDNA clone INAGE:897339 5' similar to gb:M22382 MITOCHONDRIAL MATRIX PROTEIN P1 PRECURSOR (HUMAN);
1839	11736	21611	1:11	6.8E-02		EST_HUMAN	ae30f02.r1 Gessler Wilms tumor Homo saplens cDNA clons IMAGE:897339 5' similar to gb:M22382 MITOCHONDRIAL MATRIX PROTEIN P1 PRECURSOR (HUMAN);
1865	11761	21635	3.91	6.8E-02		N	Homo saplens putative hepatic transcription factor (WBSCR14) gene, complete ods
3062	12989		1.23	6.8E-02	Γ	EST_HUMAN	ai75a06.s1 Soares_testis_NHT Homo saplens ci2NA clone 1376626 3'
3062	12989	22781	1.23	6.8E-02	6.8E-02 AA781996.1	EST_HUMAN	ai75a06.s1 Soares_testis_NHT Homo sapiens clONA clone 1376626 3'
3062	12989		1.23	6.8E-02	6.8E-02 AA781996.1	EST_HUMAN	ai75a06.s1 Soares_testis_NHT Homo saplens dJNA clone 1376626 3'
4453	14347		0.82	6.8E-02	6.8E-02 BE141076.1	EST_HUMAN	MR0-HT0069-071099-001-c05 HT0069 Homo sapiens cDNA
6315	16178		17.71	6.8E-02		LN	Homo sapiens chromosome 21 segment HS21C()68
6822	16701	26894	6.12	6.8E-02	6.8E-02 AJ248287.1	NT	Pyrococcus abyssi complete genome; segment 5/6
6822	16701	26895	6.12	6.8E-02		NT	Pyrococcus abyssi complete genome; segment 5/6
8011	19719			6.8E-02	T03214.1	EST_HUMAN	FB4A8 Fetal brain, Stratagene Homo saplens cDNA clone FB4A8 3'end similar to LINE-1
9140	18895		2.98	6.8E-02	6.8E-02 AA758014.1	EST_HUMAN	ah67f05.s1 Soares_testis_NHT Homo sapiens cDNA clone 13207053'
9764	19284		2.97	6.8E-02	9910585 NT	NT	Mus musculus latent TGF beta binding protein (Tgfb), mRNA
1511	11416		2.17	8.7E-02	AF115538.1	NT	Oncorhynchus mykiss TAP1 protein (OnmyTAP1) mRNA, OnmyTAP1*01 allele, complete cds
1851	11747	21622	5.5		A1220285.1	EST HUMAN	qg79e04.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:1841406 3'
3656	13570	23356			P17278	SWISSPROT	HOMEOBOX PROTEIN HOX-D4 (CHOX-A)
1326	11233	21089	1.05		6.6E-02 AI735509.1	EST_HUMAN	aft2e09.x1 Barstead aorta HPLRB6 Homo sapien's cDNA clone IMAGE:2354920 3' similar to SW:LIN1_NYCCO P08548 LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.;
1347	11253					Į.	Drosophila melanogaster cactin mRNA, complete cds
2133	12021	21918	3.07	8.6E-02		IN	Mus musculus Capn12 gene for calpain 12, exons 1-21, three alternative transcipts
3133	13058		1.32	6.6E-02	Q13585	SWISSPROT	MELATONIN-RELATED RECEPTOR (H9)
3418	13335	23139	8.61	6.6E-02	R64306.1	EST_HUMAN	y18b10.s1 Scares placenta Nb2HP Homo capiens cDNA clone IMAGE:139579 3'
3432	13349		2.19	6.6E-02	7108357 NT	NT	Homo sepiens mesothelin (MSLN), transcript variant 1, mRNA
3432	13349	23155	2.19	6.6E-02	7108357 NT	IN	Homo sepiens mesothelin (MSLN), transcript variant 1, mRNA
3989	13896				AF260225.1	LN.	Homo sepiens TESTIN 2 and TESTIN 3 genes, complete cds, alternatively spliced
4904	14784	24558	8.4	١	Q61703	SWISSPROT	INTER-ALPHA-TRYPSIN INHIBITOR HEAVY CHAIN HZ PRECURSOR (ITI HEAVY CHAIN HZ)
4904	14784			6.6E-02.C	Q61703	SWISSPROT	INTER-ALPHA-IRYPSIN INHIBITOR HEAVY CHAIN HZ PRECORSOR (III HEAVY CHAIN HZ)

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_			_		_	_	-	_	_		_	_	_	_			_	-	_		_				\neg				
	Top Hit Descriptor	Amsacta albistriga nucleopolyhedrovirus AcORF 17 homolog gene, complete cds	Vibrio cholerae chromosome II, section 2 of 93 of the complete chromosome	P.vulgaris mRNA for chalcone synthase	qh41d01.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:1847233 3'	Homo saplens chemokina receptor CXCR4 gene promoter region and complete cds	Homo sapiens EWS, gar22, rrp22 and bam22 genes	MR1-SN0064-010600-006-a12 SN0064 Homo sapiens cDNA	Mus musculus DIPB gene (Dipb), mRNA	Rattus norvegicus cytochrome P450 2E1 (CYP2E:1) gene, 5' flanking region	601671046F1 NIH_MGC_20 Homo sapiens cDNA clone IMAGE:3954178 6'	Homo sapiens E2F-like protein (LOC51270), mRNA	Xenopus laevis alpha(E)-catenin mRNA, complete cds	Aquifex aeolicus section 96 of 109 of the complete genome	z48h12.s1 Soares ovary tumor NbHOT Homo scpiens cDNA clone IMAGE:756743 3' similar to gb:M26038 HLA CLASS II HISTOCOMPATIBILITY ANTIGEN, DR-5 BETA CHAIN (HUMAN);	z32g05.s1 Soares_NhHMPu_S1 Homo sapiens cDNA clone IMAGE:665144 3'	Rabbit microsomal epoxide hydrolase	Nectria haematococca kinesin related protein 2 (KRP2) gene, complete cds	A.carterae precursor of peridinin-chlorophylla-protein (PCP) gene	Mus musculus histone deacetylase 5 (Hdac5), mRNA	Mus musculus histone deacetylase 5 (Hdac5), mRNA	qe07b01.x1 Soares_testis_NHT Homo seplens cDNA clone IMAGE:1738249 3' similar to contains LTR8.b3 LTR8 repetitive element;	Helerodera glycines beta-1,4-endogiucanase-1 précursor (HG-eng-1) gene, complete cds	Heterodera glycines beta-1,4-endoglucanase-1 precursor (HG-eng-1) gene, complete cds	601680425R2 NIH_MGC_83 Homo saplens cDNA clone IMAGE:3950503 3'	Mus musculus chaperonin subunit 6a (zeta) (Cct6s), mRNA	k1419.seq.F Human fetai heart, Lambda ZAP Express Homo sapiens cDNA 5'	Homo saplens mRNA for KIAA0554 protein, partial cds	Human hereditary haemochromatosis region, histone 2A-like protein gene, hereditary haemochromatosis (HLA-H) gene, RoRet gene, and sodium phosphate transporter (NPT3) gene, complete cds
	Top Hit Database Source	TN	NT	NT	EST_HUMAN	NT	IN	EST_HUMAN	TN	. LN	EST_HUMAN	LN	TN	TN	EST HUMAN	EST_HUMAN	۲	TN	Ŋ	IN	NT	EST_HUMAN	N	N	EST_HUMAN	N	EST_HUMAN	LN	LN TN
	Top Hit Acession No.	AF204882.1	AE004345.1	X08411.1	AI243326.1	AF052572.1	Y07848.1	BF374248.1	10 10 10 10 10 10 10 10 10 10 10 10 10 1	AF167430.1	BF027639.1	7706068 NT	U47624.1	AE000764.1	AA443991.1	AA195648.1	M21496.1	AF102993.1	X94549.1	6996923 NT	6986923 NT	AI191956.1	AF052733.1	AF052733.1	BE97448.1	6753323 NT	AA093305.1	AB011126.1	U91328.1
	Most Similar (Top) Hit BLAST E Value						6.6E-02		6.6E-02	6.6E-02	6.5E-02	6.5E-02		6.6E-02	6.5E-02		6.5E-02			6.4E-02	6.4E-02			6.4E-02		6.4E-02			6.4E-02
	Expression Signal	1.34	0.84	3.09	2.83	1.48	1.37	6.28	2.08	1.46	1.91	1.75	4.17	2.16	1.76	5.61	3.63	3.73	1.53	1.35	8.78	1.4	7.58	7.58	5.23	2.57	4.12	202	1.86
	ORF SEQ ID NO:	24800	24817	26998	20105	26706	27894	28457			20313	20743	21131		25383				20307			25094	L					27668	
	Exon SEQ ID NO:	15033	15053	15873	15969	16515	17656	18207	19175	19366	10508	10895	11275	11603	15333	1	18823	18051	10501	12909	15044	15267	_	! _	L	<u> </u>		<u> </u>	18754
	Probe SEQ ID NO:	5167	5190	5968	6209	6635	7806	8330	9593	9882	287	971	1369	1702	5413	8035	9034	9393	561	4802	5180	5346	5718	5718	6029	6853	7034	7603	8946

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Probe SEQ ID NO:	Exen SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acesslon No.	Top Hit Database Source	Top Hit Descriptor
8946	18754	29050	1.86	6.4E-02	U91328.1	TN	Human hereditary haemochromatosis region, histone 2A-like protein gene, hereditary haemochromatosis (HLA-H) gene, RoRet gene, and sodium phosphate transporter (NPT3) gene, complete cds
828	19635		3.65	6.4E-02	AF107890.1	TN	Homo saplens mucin 5B (MUC5B) gene, partial cds
9337	19017	25295	2.27	6.4E-02	AJ277174.1	TN	Drosophija melanogaster mRNA for mod(mdg4)51.4 protein
1720	11621	21490	. 2.	6.3E-02	AF109905.1	. L N	Mus musculus major histocompatibility locus claqs III regions Hsc70t gene, partial cds; smRNP, G7A, NG23, MutS homolog, QLCP, NG24, NG25, and NG26 genes, complete cds; and unknown genes
3552	1		2.09		P37092	SWISSPROT	HEAT SHOCK PROTEIN 70 HOMOLOG
7762	17602	27825	3.14		AB010162.1	NT	Hepatitis G virus RNA for polyprotein (NS5A region), partial cds, strain: CMR-152
8038	16637	25741	3.29	6.3E-02	BF210736.1	EST_HUMAN	601873316F1 NIH_MGC_54 Homo sapiens cDNA clone IMAGE:4097499 5'
8373	19039		1.49		P15276	SWISSPROT	TRANSCRIPTIONAL REGULATORY PROTEIN ALGP (ALGINATE REGULATORY PROTEIN ALGR3)
4167	14057	23831	3.37	6.2E-02	AL161572.2	LN LN	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 68
							Rattus norvegicus differentation-associated Na-dependent inorganic phosphate cotransporter (DNPI) mRNA,
4243	14142		1.11		AF271235.1	NT	complete cds
4479			5.41	6.2E-02	Q62191	SWISSPROT	52 KD RO PROTEIN (SJOGREN SYNDROME TYPE A ANTIGEN (SS-A)) (RO(SS-A)) (RO52)
4803	14687		1.22		AV70570	EST_HUMAN	AV705701 ADB Homo saplens cDNA clone ADBBAB03 5'
7459	17319		1.21		TN 88877898	F	Mus musculus stromal cell derived factor receptor 2 (Sdfr2), mRNA
8655	18544	28827	1.84		AJZ42735.1	NT	Melarhizium anisopliae mRNA for Chymotrypsin (chyf gene)
9129	19752		3.63		AE000750.1	LN	Aquifex aedicus section 82 of 109 of the complete genome
9541	10142	25285	1 98	6.7E-02	BF112039 1	FST HIMAN	7/37h08.x1 Sogres_NSF_F9_9W_OT_PA_P_S1 Homo sapiens cDNA clone IMAGE:3523815 3' similar to TR:0974S6 09Y4S6 HYPOTHETICAL 30.3 KD PROTEIN. [1] :
258	L	_	4.63		D16471.1	N	Human mRNA, Xq terminal portion
3909	13818		2.65		U73325.1	TN	Arabidopsis thaliana K+ inward rectifying channel protein (AtKC1) gene, complete cds
5150	15017		1.01		AB040897.1	TN	Homo sapiens mRNA for KIAA1464 protein, partitil cds
6809	16688	26877	3.48	L	X99268.1	TN	H.saplens mRNA for B-HLH DNA binding protein
8112	18002	28248	5.44	_	BE179543.1	EST_HUMAN	L3-HT0618-110500-138-C06 HT0618 Homo sapiens oDNA
8088			71.17	L	X70969.1	N	S Japanicum mRNA for serine-enzyme
9783	19286		3.6	6.1E-02	AL163207.2	N-	Homo sapiens chromosome 21 segment HS21C007
1241	11148	20997	1.41	6.0E-02	AE001777.1	TN	Thermotoga maritima section 89 of 136 of the complete genome
2641	12508	L	1.57	6.0E-02	AW968848.1	EST_HUMAN	EST380924 MAGE resequences, MAGJ Homo sapiens cDNA
2745	12607		1.61	6.0E-02	AB031289.1	TN	Mesocestoides corti mitochondrial DNA, NADH déhydrogenase subunit 4, tRNA-Gin, tRNA-Phe, tRNA-Met, ATPase subunit 6, and NADH dehydrogenase subunit 2
2906) J	19892	1.22	6.0E-02	AA188730.1	EST HUMAN	zp78c04.r1 Stratagene Hel a cell s3 937216 Homo sapiens cDNA clone IMAGE:626310 5

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
2906	10076	19893	1.22	6.0E-02	AA188730.1	EST_HUMAN	zp78c04.r1 Stratagene HeLa cell s3 937216 Homo sapiens cDNA clone IMAGE:626310 5'
3191	13116		1.2	6.0E-02	AA372376.1	EST_HUMAN	EST84288 Colon adenocarcinoma.IV Homo sapiens cDNA 5' end similar to tissue-specific protein
3191	13116	22922	1.2	6.0E-02	AA372376.1	EST_HUMAN	EST84266 Cdon adenocarcinoma IV Homo sapiens cDNA 5' end similar to tissue-specific protein
3583	13497		6.0	6.0E-02	BE984443.2	EST_HUMAN	601658150R1 NIH_MGC_68 Homo sapiens cDNA clone IMAGE:3876060 3'
4883	14773	24551	1.17	6.0E-02	267739.2	NT	Streptococcus pneumoniae parC, parE and transposase genes and ORF DNA
5314	15235		3.46		AW370211.1	EST_HUMAN	RC3-BT0253-011199-013-b04 BT0253 Homo seplens cDNA
6172	15129	24848	2.86	6.0E-02	- 5174698 NT	TN	Homo sapiens stimulated trans-acting factor (50 kDa) (STAF50) mRNA
6172	15129	24849	2.86		5174698	NT	Homo sapiens stimulated trans-acting factor (50 kDa) (STAF50) mRNA
6265	16130	26284	1.97	6.0E-02	BF382349.1	EST_HUMAN	601815274F2 NIH_MGC_56 Hamo saplens cDNA clone IMAGE:4049226 5'
6524	16383	26562	2.05	6.0E-02	AI204275.1	EST_HUMAN	qf58b08.xf Soares_testts_NHT Homo sapiens cDNA clone IMAGE:17541993'
7340	17208	27408	1.19	6.0E-02	Al623167.1	EST_HUMAN	ts78a08.x1 NCI_CGAP_GC8 Homo saptens cDNA clone IMAGE:2237362 3'
7340	17208	27407	1.19	8.0E-02	AI623167.1	EST_HUMAN	ts78a06.x1 NCI_CGAP_GC8 Homo saptens cDNA clone IMAGE:2237362 3'
7411	17278	27486	1.79	6.0E-02	AJ245365.1	NT	Acipenser baeri partial IGLV gene for Immunoglobulin light chain variable region, exons 1-2
7411	17278		1.79	6.0E-02	AJ245365.1	TN	Acipenser baeri partial IGLV gene for Immunoglobulin light chain variable region, exons 1-2
9336	19016		1.95	6.0E-02	11431702 NT	LN L	Homo sapiens DNA-dependent protein kinase catalytic subunit-interacting protein 2 (KIP2), mRNA
							wf69h03.x1 Soares_NFL_T_GBC_S1 Homo sapiens oDNA clone IMAGE:2360885 3' similar to TR:060298
9715	19258		1.84	6.0E-02	AI809273.1	EST_HUMAN	O60298 KIAA0551 PROTEIN;
229	10188	20011	3.42	5.9E-02	AW634719.1	EST_HUMAN	RC1-DT0001-290100-012-e10 DT0001 Homo suplens cDNA
2955		22681	2.59	5.9E-02	AF190269.1	IN	Mus musculus p53 tumor suppressor gene, exon 10 and 11, partial ods; alternatively spliced
4770	L		0.88	5.9E-02	AF006304.1	TN	Saccharomyces cerevisiae protein tyrosine phosphatase (PTP3) gene, complete cds
7008	16885	27077	1.87	5.9E-02	9055249 NT	NT	Mus musculus iroquols related homeobox 5 (Drosophila) (Inx5), mRNA
8165	18053		2.72	5.9E-02		NT	Mus musculus follistatin-like (Fett), mRNA
8383	18260	28509	3.15	5.9E-02	11433356 NT	NT	Homo sapiens ninein (LOC51199), mRNA
8842	18655		1.99	5.9E-02	AJ240733.1.	NT	Gallus gallus HKC9 telomere junction
917	10841		4.35	5.8E-02	D90110.1	NT	Thiobacillus ferrooxidans merC, merA genes and URF-1
2830	12759		1.1		AJ223621.1	NT	Populus trichocarpa CCoAOMT1 gene, exon 1 to exon 5
3613	13527	23314	1.44	5.8E-02	AE001775.1	NT	Thermotoga martitms section 87 of 138 of the complete genome
4257	14156		4.36		AW051927.1	EST_HUMAN	wx24c02.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2544578 3'
4257	14156		4.36	5.8E-02	AW051927.1	EST_HUMAN	wx24c02.x1 NCI_CGAP_Kid11 Homo sepiens cDNA clone IMAGE:2544578 3'
4447	14341	24132	4.21	5.8E-02	AI247505.1	EST_HUMAN	qh56f01.x1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:1848697 3' similar to gb:M13142 COAGULATION FACTOR XI PRECURSOR (HUMAN);
4447	14341	24133	4.21	5.8E-02	AI247505.1	EST_HUMAN	qh56f01.x1 Soares_fetal_liver_epleen_1NFLS_S1 Homo sapiens oDNA clone IMAGE:1848697 3' similar to gb:M13142 COAGULATION FACTOR XI PRECURSOR (HUMAN);
4471	li		2.04	5.8E-02	AF096264.1	NT	Gallus gallus tyrosine kinase JAK1 (JAK1) mRNAi, complete cds

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Single Lyones Lybressed arreadile	Top Hit Descriptor	Mus musculus tuftelin 1 (Tuft1), mRNA	Homo sapiens el F4E-transporter (4E-T), mRNA	Homo sapiens elF4E-transporter (4E-T), mRNA	Mus musculus second IL11 receptor alpha chain (IL11Ra2) gene, exons 1 and 2	Citrobacter freundii DSM 30040 cyclopropane fatty ecid synthase (cfa) gene, partial cds, dihydroxyacetone kinase (dhaK), civcerol dehydrocenase (dhaD), transcribtional ectivator (dhaR), d.3-propanediol	dehydrogenese (dhaT), giyoerol dehydratase (dhaB).>	Oryza sativa rbbi3-1 gene for putative Bowman Birk trypsin inhibitor	RC5-BT0559-140200-012-C03 BT0559 Homo saplens cDNA	Xenopus laevis homeobox protein (Vox-1) mRNA, complete cds	Mus musculus p-glycoprotein (mdrla) gene, excins 1 and 2	Neurospora crassa ubiquinol-cytochrome c oxidoreductase subunit VIII (QCR8) mRNA, complete cds	Rana catesbiana heat shock protein 30 (HSP30) mRNA, complete cds	QV0-ST0213-021299-062-e09 ST0213 Homo sapiens cDNA	QV0-ST0213-021299-062-a09 ST0213 Homo sapiens cDNA	y837H2.r1 Stratagene lung (#937210) Homo saplens cDNA clone IMAGE:119951 5' similar to gb:K01506 HLA CLASS II HISTOCOMPATIBILITY ANTICEN, DP(1) ALPHA CHAIN (HUMAN);	Pseudomonas putida ttgS gene	Drosophila melanogaster laminin B2 gene, complete cds	Drosophila melanogaster laminin B2 gene, complete cds	Pseudomonas putida ttgS gene	Arabidopsis thallana eli5 gene, exons 1-11	Mus musculus caudal type homeobox-1 (Cdx-1) gene, complete cds	Helicobacter pylori 26695 section 5 of 134 of the complete genome	Helicobacter pylori 26695 section 5 of 134 of the complete genome	Lymphocystis disease virus 1, complete genome	nuclear protein TIF1 isoform [mice, mRNA, 4055 nt]	Podospora anserina mitochondrial epsilon-sen DNA	Homo sapiens meprin A, alpha (PABA peptide hydrolase) (MEP1A) mRNA	Homo sapiens partial LMO1 gene for LIM domali, only 1 protein, exon 1	Homo sepiens partial LMO1 gene for LIM domain only 1 protein, exon 1	Human steroid hormone receptor Ner-I mRNA, complete cds	Drosophila melanogaster filament protein homolog (sep1) gene, complete cds
ום ראסוו ביוסט	Top Hit Database Source				NT.	<u> </u>	Z		EST_HUMAN	NT	I.	TN	Z	EST_HUMAN (EST_HUMAN (EST_HUMAN	Г			IN		LN	INT	NT		TN	NT					LN LN
8110	Top Hit Acesslon No.	6755902 NT	10947034 NT	10947034 NT	U69492.1		U09771.1	-	BE073468.1	U53528.1	M96761.1			18.1	AW391248.1					•	5.3E-02 AJ011048.1		-	.1	9695413 NT			5031908 NT		.1		
	Most Similar (Top) Hit BLAST E Value	5.5E-02	5.5E-02	5.6E-02			5.5E-02		5.4E-02		5.4E-02	5.4E-02 U20790.1	5.4E-02 U44894.1	5.3E-02	5.3E-02	5.3E-02 T94759.1	5.3E-02	5.3E-02 M58417.1	5.3E-02 M58417.1	5.3E-02	5.3E-02	5.3E-02	5.3E-02 AE000527	5.3E-02	5.3E-02	5.3E-02 S78221.1	5.3E-02 X03127.1	5.2E-02	5.2E-02	5.2E-02	5.2E-02 U07132.1	5.2E-02 L33246.1
	Expression Signal	1.58	1.3	1.3	1.48		11.56	0.93	6.34	0.92	1	1.79	1.55	1.62	1.62	18.21	3.14	3.91	3.91	4.59	1.22	7.26	1.78	1.76	3.87	1.94	1.78	439.66	2.34	2.34		1.04
	ORF SEQ ID NO:		L	27633			28508			24563		28223		20797						L		24662					27349		22793		23854	L
	Exan SEQ ID NO:	16228	17417	1	1		18259	I_	15078	14787	14976	L.	1	10955	10955		12324	_		13038	14399	14894	1	1	Į	١	17153	12123	13003	13003	14081	14521
	Probe SEQ ID NO:	6365	7568	7566	7619		8382	2986	3375	4908	5108	8083	8323	1037	1037	1489	2447	2912	2912	3113	4508	5021	6268	5258	6115	6353	7276	2239	3076	3076	4181	4633

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Top Hit Descriptor	wj80e04.xt NCI_CGAP_Lym12 Homo sepiens cDNA clone IMAGE:2409150 3' similar to contains MER15.b1 MER15 repetitive element ;	Homo sapiens chromosome 21 segment HS21C004	Turnip mosaic virus genomic RNA for Capsid protein, complete cds	Turnip mosaic virus genomic RNA for Capsid protein, complete cds	OXALOACETATE DECARBOXYLASE ALPHA CHAIN	DKFZp547D073_r1 547 (synonym: hfbr1) Horrio sapiens cDNA clone DKFZp547D073 5'	601653565R2 NIH_MGC_55 Homo sapiens cDNA clone IMAGE:3838361 3'	QV0-UM0051-250800-350-b08 UM0051 Homp saplens cDNA	Spodoptera littoralis mRNA for 3-dehydroecdysone 3beta-reductase	Candida albicans protein phosphatase Ssd1 hcmolog (SSD1) gene, complete cds	ANTER-SPECIFIC PROLINE-RICH PROTEIN APG (PROTEIN CEX)	Homo saplens ES18 mRNA, partial cds	Homo saplens ES18 mRNA, partial cds	Cucumis melo polygalacturonase precursor (MPG3) mRNA, complete cds	Mus musculus fatty acid amide hydrolase gene, exon 10	Bacillus subtilis complete genome (section 1 of 21): from 1 to 213080	SALIVARY ACIDIC PROLINE-RICH PHOSPHOPROTEIN 1/2 PRECURSOR (PRP-1/PRP-3) (PRP-2/PRP- 4) (PIF-FIPIF-S) (PROTEIN A/PROTEIN C) [CONTAINS: PEPTIDE P-C]	Oryctolegus cuniculus UDP-glucuronosyltransferase (UGT2B13) mRNA, complete cds	Mus musculus Unc-51 like kinase 2 (C. elegans) (UIk2), mRNA	Antheraea pernyl period clock protein homolog mRNA, complete cds	Homo sapiens ubiquitous tetratricopeptide containing protein RoXaN mRNA, partial cds	NEUROFILAMENT TRIPLET L PROTEIN (NEUROFILAMENT LIGHT POLYPEPTIDE) (NF-L)	Mus musculus Fas-interacting serine/threonine kinase 3 (Fist3) mRNA, complete cds	Methanococcus jannaschii section 142 of 150 of the complete genome	NO-ON-TRANSIENT A PROTEIN	Ohicken 28-kDa vitamin D-dependent calclum-binding protein (CaBP-28) mRNA, complete cds	Homo sapiens ABCA1 (ABCA1) gene, complete cds	Homo sapiens ABCA1 (ABCA1) gene, complete cds	ATROPHIN-1 (DENTATORUBRAL-PALLIDOLUYSIAN ATROPHY PROTEIN)	zq48a12.s1 Strategene hNT neuron (#937233) Homo sapiens cDNA clone IMAGE:632926 3' similar to contains Alu repetitive element,contains element MSR1 repetitive element;	zi78a03.s1 Soares_bstis_NHT Homo saplens cDNA clone IMAGE:728428 3'
Top Hit Database Source	EST_HUMAN	LN.	NT	NT	SWISSPROT	EST_HUMAN	EST HUMAN	EST_HUMAN	NT	N	SWISSPROT	N	LN	NT	ΙΝ	L'A	SWISSPROT	N	12	N	NT	SWISSPROT	INT	NT	SWISSPROT	TN.	TN	INT	SWISSPROT	EST_HUMAN	EST_HUMAN
Top Hit Acession No.	A1830965.1	2 AL163204.2	2 D10927.1	D10927.1	2 003030	5.1E-02 AL134071.1	3E857423.2	6.1E-02 BF378625.1	5.1E-02 AJ131966.1	5.1E-02 AF012898.1	540603	AF083930.1	AF083930.1	5.1E-02 AF062467.1	5.0E-02 AF098004.1	299104.1	2 P02810	U72742.1	7305610 NT	5.0E-02 U12769.2	5.0E-02 AF188530.1	P35616	3.1	-		4.9E-02 M14230.1	4.9E-02 AF275948.1	AF275948.1	4.9E-02 P54258	AA188940.1	4.8E-02 AA400914.1
Most Similar (Top) Hit BLAST E Value	5.2E-02	5.2E-02	5.2E-02	5.2E-02	5.2E-02	5.1E-02	5.1E-02	6.1E-02				5.1E-02	5.1E-02	5.1E-02	5.0E-02	5.0E-02 Z89104.1	5.0E-02	5.0E-02 U72742.	5.0E-02									4.9E-02	4.8E-02		
Expression Signal	1.73	2.23	2.03	2.03	1.63	1.02	1.12	1.65	1.43					1.81	1.98	7.11	4.27						1.32			28.95		2.57			
ORF SEQ ID NO:			27877	27878	L		24820	24871	26930						20234			20742	L	23330							20143		22972		23262
Exen SEQ ID NO:	15822	16647	17461	17461	19167	12194	14854	15108	16738	i)	18088	18088	i _	10418	L	i	L	i .	13543	l	١	17708	L		10188	10321	1	1	1	1 1
Probe SEQ ID NO:	5714	6768	7610	7810	9561	2313	4979	8609	6889	7845	7849	8204	8204	9569	474	1186	1947	2788	3285	3629	4850	6450	7858	8782	9606	218	363	365	3251	3522	3543

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	Exan NO: NO: 13459 14618 14618 16220 18523 19107 10287 10287 10287 10287 10287 10287 10484 1484 16810 16830	유	Signa			Top Hit Database Source Source EST_HUMAN EST_HUMAN NT NT NT NT NT NT NT NT NT EST_HUMAN NT NT EST_HUMAN NT EST_HUMAN NT EST_HUMAN NT EST_HUMAN NT EST_HUMAN NT EST_HUMAN NT EST_HUMAN NT EST_HUMAN NT EST_HUMAN NT EST_HUMAN NT NT EST_HUMAN NT EST_HUMAN NT NT EST_HUMAN NT EST_HUMAN NT NT EST_HUMAN NT NT NT NT NT NT NT NT NT NT NT NT NT	Top Hit Descriptor Top Hi
8800 8800	15981				M62752.1 X15543.1	Z	Rati Batin-related protein (s.) yetre, compress c.c.o. Malarus mRNA for RF-36-DNA-binding protein. U socioso DNA for and communic retroined like element
7179	17056			4.7E-02 4.7E-02	X89211.1 AB026678.1 X46643.4	12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 1	H.seplens DNA for endogenous retroviral like element Gallus gallus Wpkci-8 gene, complete cds R Jaunis mRNA for RF-36-DNA-binding protein
7321 8136 9305	18024 19740	27397 28270		4.7E-02 4.7E-02 4.7E-02	X15543.1 8754565 NT AV648521.1 ES	NT NT EST_HUMAN	B. taurus mknya ter kr3e-Unya-entaing protein Mus musculus ligand of numb-protein X (Lmx), mRNA AV648621 GLC Homo septens cDNA clone GLCBKD023'
9852	19743		2.35		P52951 AJ277662.1	SWISSPROT	HOMEOBOX PROTEIN GBX-2 (GASTRULATION AND BRAIN-SPECIFIC HOMEOBOX PROTEIN 2) Homo sapiens partial TUB gene for tubby (mousė) homolog and LMO1 gene for LIM domain only 1 protein
1270	1 1 1	20484			AE000445.1 Al014266.1	NT EST_HUMAN	Escherichia coli K-12 MC1655 section 335 of 400 of the complete genome am50d02.s1 Johnston frontal cortex Homo sapiens cDNA clone IMAGE:1538979 3' similar to TR:P90533 P90533 LIMA; contains element LTR1 repetitive element;

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Table 4
Single Exon Probes Expressed in Heart

			Most Similar		107	
SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	(Top) Hit BLASTE Value	Tap Hit Acesslan No.		Top Hit Descriptor
11242	21100	2.74	4.6E-02	AV727059.1	EST_HUMAN	AV727059 HTC Homo sepiens cDNA clone HTCBWC01 6
12315			4.6E-02	AW236023.1	EST HUMAN	xn24f03.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2894653 3' similar to SW:GRF1_HUMAN Q12849 G-RICH SEQUENCE FACTOR-1;
10235			4.6E-02	BE153583.1	EST_HUMAN	PM0-HT0339-251199-003-g05 HT0339 Homo capiens cDNA
12904			4.6E-02	BE153583.1	EST_HUMAN	PM0-HT0339-251199-003-g05 HT0339 Homo sapiens cDNA
12904	4 22703	0.95	4.6E-02	BE153583.1	EST_HUMAN	PM0-HT0339-251199-003-g05 HT0339 Homo caplens cDNA
13936		1.22	4.6E-02	AF220365.1	NT	Mus musculus nucleolar RNA helicase II/Gu (ddx21) gene, complete ods
15684	4 25792	3.64	4.6E-02	X61624.1	NT	C.reinhardtii atp2 (atpB) mRNA
15684			4.6E-02	X61624.1	NT	C.reinhardii atp2 (atpB) mRNA
16063	28200	1 31	4 GF-02	A1149574 1	EST HUMAN	qc60b06.x1 Soares_placenta_8tc9weeks_2NbHP9to9W Homo seplens cDNA done IMAGE:1713971 3' similar to contains L1:t3 L1 repetitive element ;
200			4 6F-02	BE154008 1	EST HUMAN	PM0-HT0339-060400-009-G12 HT0339 Homo sapiens cDNA
18524			4.6E-02	AA913328.1	EST HUMAN	ol27h09.s1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:1524737 3'
19738			4.6E-02		NT	Oryctolagus cuniculus macrophage scavenger receptor type II mRNA, complete cds
19343	3	2.68			NT	Human germline immunoglobulin lambda light chain gene
10383	3 20207	1.84	4.5E-02	P22448	SWISSPROT	RETINOIC ACID RECEPTOR BETA (RAR-BETA)
11110			4.5E-02	AF005730.1	NT	Marburg virus strain M/S. Africa/Johannesburg/1975/Ozolin VP35 gene, complete cds
11110	0 20958	0.85	4.6E-02	AF005730.1	NT	Marburg virus strain MIS. Africal Johannesburg/1975/Ozolin VP35 gene, complete cds
11862		3.55	4.5E-02	P32182	SWISSPROT	HEPATOCYTE NUCLEAR FACTOR 3-BETA (HNF-3B)
11953		2.04	4.5E-02	AE003964.1	L	Xylella fastidiosa, section 110 of 229 of the complete genome
13576		3.83	4.6E-02	AL163278.2	NT	Homo sapiens chromosome 21 segment HS21Co78
			100	7	į.	Homo sapiens ASCL3 gene, CEGP1 gene, C11orf14 gene, C11orf15 gene, C11orf16 gene and C11orf17
13083	28085	1.04	4.3E-02 4.5F-02	AJ4006/7.1	- L	Arabidopsis thallana CCAAT-box binding factor HAP3 homolog gene, complete cds
17568			4.5E-02		EST HUMAN	EST28167 Cerebellum II Homo sapiens cDNA 5' end similar to similar to neuro-D4 protein
19000			4.5E-02		TN	Homo saplens ret finger protein-like 3 (RFPL3), mRNA
19637				AA191097.1	EST_HUMAN	zq43f11.r1 Stratagene hNT neuron (#937233) Homo sapiens cDNA clone IMAGE:632493 5
10184	L		1	BE972733.1	EST_HUMAN	601652154F1 NIH_MGC_82 Homo sapiens cDNA clone IMAGE:3935388 5'
10926	5 20770	1.29	4.4E-02	L19295.1	NT	Drosophila melanogaster extradenticle (EXD) mRNA, complete cds
11940	0	6.15	}	P31568	SWISSPROT	HYPOTHETICAL PROTEIN (ORF 2280)
12317	7 22214	1.29	4.4E-02	AW875475.1	EST_HUMAN	QV2-PT0012-010300-070-g02 PT0012 Homo stiplens cDNA
13502	12 23291	1.88	4.4E-02	AF159160.1	본	Myxococcus xanthus serine/threonine kinase Pkr10 (pkn10) gene, complete cds
14420		1.08	4.4E-02	AF109907.1	NT	Homo sapiens S164 gene, pariial cds; PS1 and hypothetical protein genes, complete cds; and S171 gene, partial cds
1	١					

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Table 4
Single Exon Probes Expressed in Heart

Probe SEQ ID NO:	Exen SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
4527	14420	24204	1.08	4.4E-02	AF109907.1	IVI	Homo sepiens S164 gene, partial cds; PS1 and hypothetical protein genes, complete cds; and S171 gene, partial cds
4637	14525		3.12	4.4E-02	AJ222689.1	LN T	Ovis aries CCAAT-enhancer binding protein epsilon gene
7084	16961	27154	1.98	4.4E-02	AA736969.1	EST_HUMAN	nw13h03.81 NCI_CGAP_SS1 Homo septens cDNA clone IMAGE:1239221 3'
8423	18297	28552	4.11	4.4E-02	AF060669.1	۲N	Hepatitis E virus strain HEV-JUS2 polyprotein (ORF1), (ORF3), and capsid protein (ORF2) genes, complete cds
8533		28670	2.39	4.4E-02	AA496739.1	EST_HUMAN	ae33f04.r1 Gessler Wilms turnor Homo sapiens cDNA clone IMAGE:897631 5'
8028	18820		1.88	4.4E-02	AB040926.1	F	Homo sapiens mRNA for KIAA1493 protein, partial cds
9210	19753		1.44	4.4E-02	BF241245.1	T_HUMAN	601878746F1 NIH_MGC_55 Homo sepiens cDNA clone IMAGE:4107418 5
784	10695		5.74	4.3E-02	AF003249.1	TN	Morone saxatilis myosin heavy chain FM3A (FM3A) mRNA, complete cds
2521	12395	22286	1.23	_	AV704878.1	T_HUMAN	AV704878 ADB Homo sapiens cDNA clone ADBAOH08 5'
3383	13301	23101	7.04	4.3E-02	AL163210.2		Homo sapiens chromosome 21 segment HS21C010
3609	13523		1.07	4.3E-02		ΤN	Homo saplens promyelocytic leukemia zinc finger protein (PLZF) gene, complete cds
5141	16008	24779	0.92	4.3E-02	X51594.1	TN	Pea P4 organ specific gene
5914	15820	25944	4.93	4.3E-02	P30427	SWISSPROT	PLECTIN
5914	15820	25945	4.93	4.3E-02	P30427	SWISSPROT	PLECTIN
8242		28372	2.48	4.3E-02	X17012.1	LN	Rat IGFII gene for insulin-like growth factor II
805	10734	20577	2.05		AU123327.1	EST_HUMAN	AU123327 NT2RM2 Homo sapiens cDNA clone NT2RM2000020 5'
848	10775		1.93	4.2E-02	AU123327.1	EST_HUMAN	AU123327 NT2RM2 Homo sapiens cDNA done NT2RM2000020 6
878	10804	20654	0.83	4.2E-02	AW003645.1	EST HUMAN	wo34g01.x1 NCI_CGAP_Pit1 Homo saplens cDNA clone IMAGE:2545584 3' similar to TR:Q63291 Q63291 L1 RETROPOSON, ORF2 MRNA contains L1.t3 L1 L1 repetitive element:
1690	11592		1.21	_	AL445066.1	NT	Thermoplasma acidophilum complete ganome; sagment 4/5
3117	13042	22838	0.8	4.2E-02)	A1493472.1	EST_HUMAN	qy95f10.x1 NCI_CGAP_Brn25 Homo sapiens cDIVA clone IMAGE:2019787 3' similar to gb:M35718 FIBROBLAST GROWTH FACTOR RECEPTOR BFR-2 PRECURSOR (HUMAN);
5172	15038		1.07	4.2E-02		NT	Human mRNA for KIAA0150 gene, partial cds
6444	16305	28470	4.45		2.1	IN	Legionella pneumophila catalase-perceidase (katA) gene, complete cds
7114	16991	27183	3.88	4.2E-02	P05095	SWISSPROT	ALPHA-ACTININ 3, NON MUSCULAR (F-ACTIN CROSS LINKING PROTEIN)
7845	17695	27941	1.28		Q16650	SWISSPROT	T-BRAIN-1 PROTEIN (T-BOX BRAIN PROTEIN (1) (TBR-1) (TES-56)
8627	18492	28764	2.33			EST_HUMAN	PM3-BN0174-250500-009-d10 BN0174 Homo capiens cDNA
8627	18492	28765	2.33			EST_HUMAN	PM3-BN0174-250500-009-d10 BN0174 Homo sapiens cDNA
8795	18609	28900	1.73	_			PRRS Isolate PRRSV36 envelope glycoprotein gene, complete cds
9563	19881		2.69	_		EST HUMAN	wt48g10.x1 NCI_CGAP_Pan1 Homo sapiens cDNA clone IMAGE:2510850 3'
2843	ŀ		0.97				Chlamydia muridarum, section 60 of 85 of the complete genome
3824	13736	23526	0.86	4.1E-02	BE297236.1	EST_HUMAN	601177907F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:3533353 5'

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		_			_	_	_	_			_		_		_								_						_	
Top Hit Descriptor	601177907F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:3533353 5'	QV1-NN0012-180400-164-f06 NN0012 Homo sapiens cDNA	Homo sepiens KIAA0867 protein (KIAA0867), mRNA	Fugu rubripes neural cell adhesion molecule L1 homolog (L1-CAM) gene, complete cds; putative protein 1	(PUT1) gene, partial cds; mitosis-specific chromosome segregation protein SMC1 homolog (SMC1) gene,	complete cds; and calcium channel alpha-1 subunit>	Brassica napus gin gene for plastid glutamine syithetase, exons 1-12	Homo sapiens mRNA for KIAA1471 protein, partial cds	Homo sapiens cytochrome P450 polypeptide 43 (CYP3A43) gene, partial cds; cytochrome P450 polypeptide	4 (CYP3A4) and cytochrome P450 polypeptide 7 (CYP3A7) genes, complete cds; and cytochrome P450	polypeptide 5 (CYP3A5) gene, partial cds	Strongylocentrotus purpuratus homolog of human bone morphogenetic protein 1 (submp) mRNA, complete cds	GLUCOAMYLASE S1/S2 PRECURSOR (GLUCAN 1,4-ALPHA-GLUCOSIDASE) (1,4-ALPHA-D-GLUCAN	GLUCOHYDROLASE)	Methanobacterium thermoautotrophicum strain Marburg, Thiol:fumarate reductase subunit A	Kluyveromyces lactis gene for Ca++ ATPase	Ovis aries mRNA for acetyl-coA carboxylase	UI-H-BW1-anx-h-08-0-UI.s1 NCI_CGAP_Sub7 Homo sapiens cDNA clone IMAGE:3084134 3'	FAS ANTIGEN LIGAND	M.musculus DNA for desmin-binding fragment DesD7	Homo expiens succinate dehydrogenase complex, subunit C, integral membrane protein, 15kD (SDHC) mRNA	Homo sepiens hypothetical protein PRO1163 (PRO1163), mRNA	Homo sapiens hypothetical protein PRO1163 (PRO1163), mRNA	601906848F1 NIH_MGC_54 Homo sapiens cDNA done IMAGE:4134779 5'	Fells catus G-CSF gene for granulocyte colony-stimulating factor, complete cds	Human germline T-cell receptor beta chain TCRBV17S1A1T, TCRBV2S1, TCRBV10S1P, TCRBV29S1P,	TCRBV19S1P, TCRBV15S1, TCRBV11S1A11, HVB relic, TCRBV28S1P, TCRBV34S1, TCRBV14S1, TCRBV3S2, TCRBV3S1, TCRBV4S1A11, TRY4, TRY5, TRY7, TRY8, TCRBD1, TCRBJ1S1, TCRBJ1S2,	Mus musculus chromosome X contigB; X-linked lymphocyte regulated 5 gene, Zinc finger protein 275, Zinc	finger protein 92, mmxq28orf	601510891F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3912215 5'
Top Hit Database Source	EST_HUMAN	EST HUMAN	7662347 NT			NT	N	LV.			K	LN		SWISSPROT	Ę	F	١	EST HUMAN	SWISSPROT	N _T	LN	N-	Į,	EST HUMAN	NT		Ę		TN	EST_HUMAN
Top Hit Acession No.	BE297236.1	AW893484.1	7662347			AF026198.1	AJ271909.1	AB040904.1			AF280107.1	L23838.1		P08640	AJ000941.1	AJ001018.1	AJ001056.1	BE516149.1	P41047	AJ403386.1	TN CARACA	8924019 NT	8924019 NT	BF239613.1	AB042553.1		U66081.1			BE885137.1
Most Similar (Top) Hit BLAST E Value	4.1E-02	4.1E-02	4.1E-02			4.1E-02	4.1E-02	4.0E-02			4.0E-02	4.0E-02		4.0E-02	4.0E-02	4.0E-02	4.0E-02	3.9E-02	3.9E-02	3.9E-02	3 OF 02	3.9E-02	3.9E-02				3.9E-02		3.9E-02	3.8E-02
Expression Signal	0.88	7.37	1.84			2.44	12.83	2.68			4.92	5.69		2.8	2.42	1.94	4.98	2.77	3.48	3.04	0.00	0.98	96:0	122	5.45		1.57		7.38	0.94
ORF SEQ ID NO:	23527		26459			26604	İ	22932			25029	26567		27126	.27638		25135	20863		21691		24748				٠				21683
Exan SEQ ID NO:	13736	14268	16297			16423	19682	13130			15225	16387		16936	17423	18788	19506	11020	11229	11813	1253B			16461	19644		19248		19568	11804
Probe SEQ ID NO:	3824	4372	6438			6565	9873	3206			5304	6528		7059	7572	8983	9198	1104	1322	1918	2674	5105	5105	6581	9028		9697		9811	1909

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	Top Hit Descriptor	Homo sapiens ATP-binding cassette, sub-family A (ABC1), member 8 (ABCA8), mRNA	Human von Willebrand factor gene, exons 23 through 34	Homo sapiens PELOTA (PELOTA) gene, complete cds	LAMININ ALPHA-1 CHAIN PRECURSOR (LAMININ A CHAIN)	Homo saplens plasma membrane calclum ATP ase Isoform 1 (ATP 2B1) gene, alternative splice products, partial cds	wr85e08x1 NCI CGAP Kid11 Homo sapiens cDNA clone IMAGE:2494502 3'	Homo sapiens mRNA for KIAA0718 protein, partial cds	EOMESODERMIN	601898233F1 NIH_MGC_19 Homo sapiens cDNA done IMAGE:4126584 6	Mus musculus potassium large conductance pH·sensitive channel, subfamily M, alpha member 3 (Konma3), mRNA	601762117F1 NIH MGC_20 Homo sepiens cDNA done IMAGE:4024973 5'	Homo saplens solute carrier family 22 (organic cation transporter), member 1 (SLC22A1), mRNA	H.vulgare Ss1 gene for sucrose synthase	Homo saplens genomic region containing hypervariable minisatellites chromosome 10[10q26.3] of Homo	sepiens	Homo saplens genomic region containing hypervariable minisatellites chromosome 10[10q26.3] of Homo saplens	CM2-EN0013-110500-192-b10 EN0013 Homo sapiens cDNA	CM2-EN0013-110500-192-b10 EN0013 Homo sapiens cDNA	Chromatium vinosum sulfur globule protein Cv2 precursor (sgp2) gene, complete cds	nw20e05.s1 NCI_CGAP_GCB0 Home sepiens cDNA clone IMAGE:1241024 3' similar to gb.J00314_rna2 TUBULIN BETA-1 CHAIN (HUMAN);	Dictyostellum discoldeum unknown spore germination-specific protein-like protein, orf1, orf2 and orf3 genes,	complete cds	Dictyostalium discoldeum unknown spore germination-specific protein-like protein, orf1, orf2 and orf3 genes,	campieise cas	Drosophila metenogaster üggrin mikny, complete cas	Homo sapiens microsomal epoxide hydrolase (EPHX1) gene, complete cds	602085136F1 NIH_MGC_83 Homo sapiens cDNA clone IMAGE:4249377 5	802085136F1 NIH MGC 83 Home septens cDNA clone IMAGE:4249377 5
	Top Hit Database Source		F		SWISSPROT 1	<u> </u>	T HUMAN	Г	SWISSPROT	EST_HUMAN		T HUMAN				EN FN	L N	EST HUMAN	HUMAN	Г	EST HUMAN	Г	E					٦	DAM IN PAR
	Top Hit Acession No.	E005700 NT	M60675.1	AF143952.2	P19137	1 14561 1	-	_	P79944	BF312963.1	6680544	BF124974.1	18392	X73221.1		AL096806.1	AL096810.1		AW945516.1	AF025952.1	AA714521.1		U20608.1						DERZOOR 4
	Most Similar (Top) Hit BLAST E Value	3.8E-02	3.8E-02	3.8E-02/	3.7E-02	3 7F-02			3.7E-02		3 75 00			3.6E-02		3.6E-02	3.6E-02	3.6E-02	3.6E-02 /	3.6E-02	3.6E-02		3.6E-02		3.8E-02	3.5E-02		3.5E-02 E	2 50 00 0
	Expression Signal	1.74	1.39	2.45	4.59	å	6.3	0.91	0.8	3.45	-	3.4	1.62	0.85		0.8	0.84	5	ß	1.82	2.63		2.08		2.08	1.15	1.83	1.49	1 40
	ORF SEQ '	26365		28185		24128		22299	22733	22734		29119		23306	1	23313	24773	26062	26063	28229	26349		27473	_	1	1			24340
	Exon SEQ ID NO:	16194	16910	17938	10899	11272			L	12941	13326	L	1	13518	1	13526	15002	15932	1	i	16187		17269	_	1	_ [١		44.40
	Probe SEQ ID NO:	6331	7033	8045	976	1386	2190	2533	3012	3013	3408	9095	9755	3604		3612	5135	8028	8028	6214	6324		7402		7402	8	992	1544	1544

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	T		_	_	_		1	•	_				_	$\overline{}$	Т	Т	\neg	т	Т	T	Τ-		_	_			Т	_	Ţ
Top Hit Descriptor	CYSTATHIONINE BETA-LYASE PRECURSOR (CBL) (BETA-CYSTATHIONASE) (CYSTEINE LYASE)	EST370539 MAGE resequences, MAGE Homo saplens cDNA	HYPOTHETICAL 80.7 KD PROTEIN IN SOD1; CPA2 INTERGENIC REGION	Maize actin 1 gene (MAc1), complete cds	601644701R2 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:3929737 3'	L. lactis MG1363 grpE and dnaK genes	PM1-CT0326-291299-002-h03 CT0326 Homo sapiens cDNA	PM1-CT0326-291299-002-h03 CT0326 Homo saplens cDNA	601178765F1 NIH_MGC_20 Homo sapiens cDNA clone IMAGE;3543833 5	Homo sapiens mRNA for FLJ00013 protein, partial cds	Homo sapiens mRNA for FLJ00013 protein, partial cds	Homo sapiens mRNA for FLJ00013 protein, partial cds	Homo sapiens mRNA for FLJ00013 protein, partial cds	XZ6407.X1 Soares, NFL, T_GBC_S1 Homo sapiens cDNA clone IMAGE:2814253 3' similar to SW:C211, HUMAN P53801 PUTATIVE SURFACE GLYCOPROTEIN C210RF1 PRECURSOR;	Homo sapiens hypothetical protein FLJ13220 (FLJ13220), mRNA	yo20e08.r1 Strategene lung (#937210) Homo sepiens cDNA clone IMAGE:81250 5' similar to contains	MENZY repealuve element	RC3. ENDARS OF OUT A TANA SE HOME SERVICES ON A	RC8-UM0015-210200-021-A10 UM0015 Home sapiens cDNA	M.musculus S-antigen gene promoter region	LA PROTEIN HOMOLOG (LA RIBONUCLEOPROTEIN) (LA AUTOANTIGEN HOMOLOG)	Caenorhabditis elegans mRNA for DYS-1 protein, partial	Human lysyl oxidase-like protein gene, exon 3	wip9d04.xt NCI_CGAP_Bm25 Homo sapiens cDNA clone IMAGE:2433031 3'	zq04f11.s1 Stratagene muscle 937209 Homo saplens cDNA clone IMAGE:628749 3' similar to TR:G1017425 G1017425	IPISGKPLPKVTLSRDGVPLKATMRFNTEITAEŅLTINI.KESVTADAGRYEITAANSSGTTKAFINIVVLDRPG PPT GPVVISDITEESVTLKWEPPKYDGGSQVTNYILLKRETSTAVWTEVSATVARTMMKVMKL :	zi75e08.s1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:728198 3'	Cricetulus griseus CYP2A17 mRNA for cytochrome P450 2A17, complete cds	
Top Hit Database Source	SWISSPROT	EST_HUMAN	ISSPROT	IN	EST_HUMAN	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	·	IN	Ę	EST HUMAN			EST TOWAN	T LI INCON	Т	Г	SWISSPROT	TN		EST_HUMAN		EST HUMAN	Г		
Top Hit Acession No.	P63780	AW958469.1	P47144	J01238.1	BE958970.1	X76642.1	AW861641.1	AW861641.1	BE276948.1	AK024424.1	AK024424.1	AK024424.1	AK024424.1	AW274020.1	11345459 NT		15/160.7	ALIOSOG.	AW 794952.1	X59789.1	Q26457	AJ012469.1	U24393.1	AI869629.1		AA194306.1	AA398735.1	AB035867.1	
Most Similar (Top) Hit BLAST E Value	3.6E-02	3.5E-02	3.5E-02	3.5E-02		3.5E-02	3.5E-02	3.5E-02	3.5E-02	3.4E-02	3.4E-02	3.4E-02	3.4E-02	3.4E-02		1	3.45-02		3.4E-02	3.4E-02	3.4E-02	3.4E-02		3.4E-02		3.4E-02	3.3E-02	3.3E-02	
Expression Signal	1.43	98.0	0.84	1.88	2.35	1.72	1.76	1.78	3.51	1.78	1.78	3.31	3.31	3.77	10.22		2.00	120	3.18	2.41	3.43	1.28	4.19	4.78		6.07	13.18	13.17	
ORF SEQ ID NO:	23895			25787	27082	27829	28889	28890			20311	20310	20311	20795			22,220					24650	24868						
Exan SEQ ID NO:	14118	14429	15021	15680	16889	17605	18600	18600	19582		10504	10504	10504	10953	l	L_	12223	1	1	l_	14867	14884	15105	16687		17046	1		
Probe SEQ ID NO:	4220	4536	5154	5773	7012	7755	8785	8785	9749	564	564	595	565	1035	1188	3	200	2700	3845	4494	4992	5010	6095	8089		7169	368	1151	

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Probe SEQ ID NO:	Econ SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
1705	11606		1.28	3.3E-02	AE000700.1	TN	Aquifex aedicus section 32 of 109 of the complete genome
2038			2.77	3.3E-0	2 R09112.1	EST_HUMAN	yr25c09.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:127888-5'
4086	11525		2.24	3.3E-0	2 AF110763.1	LN TN	Homo saplens skaletal muscle LIM-protein 1 (FHL1) gene, complete cds
4366	14262	24047	1.88	3.3E-02	LN 2989929	TN	Mus musculus tumor rejection antigen gp96 (Trail), mRNA
5875	15781	25901	18.26		3.3E-02 BF245995.1	EST_HUMAN	601853910F1 NIH_MGC_67 Homo sapiens cDNA clone IMAGE:4073787 5'
5875	15781	25902	18.26	3.3E-0	2 BF245995.1	EST_HUMAN	601853910F1 NIH_MGC_57 Homo saplens cDNA clone IMAGE:4073787 5'
8464	18337		3.39	3.3E-02	2 BF691107.1	EST_HUMAN	602247171F1 NIH_MGC_62 Homo sapiens cDNA clone IMAGE:4332497 5'
9290	18991		1.78	3.3E-02		EST_HUMAN	ye49f11.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:121101 67
9441	19078		1.59	3.3E-02	3.3E-02 M81890.1	N	Human Interleukin 11 (IL11) gene, complete mRNA
127	10101		1.13	3.2E-02		M	Oryctolagus cuniculus gene encoding ileal sodium-dependent bile acid transporter
1110	11025		12.7	3.2E-02	3.2E-02 AF096275.1	Ę	Drosophila melanogaster heat shock protein 68 (hsp68) gene, hsp68d allele, complete cds
1110	11025	20868	12.7	3.2E-02	AF096275.1	NT.	Drosophila melanogaster heat shock protein 68 (hsp68) gene, hsp68d allele, complete cds
1734	11635		1.14	3.2E-02	AF128894.1	Z	Homo saplens telomerase reverse transcriptase (TERT) gene, exons 7-16 and complete cds
2072	11962		16.0	3.2E-02	2 P28955	SWISSPROT	LARGE TEGUMENT PROTEIN
2809	10101	19923	1.15	3.2€-02	AJ002005.1	NT	Oryctolagus cuniculus gene encoding ileal sodium-dependent bile acid transporter
3095	13022		10.71	3.2E-02	BE867353.1	EST_HUMAN	601442431F1 NIH_MGC_65 Homo sapiens cDNA clone IMAGE:3846727 5'
3653	13567	23354	1.05	3.2E-02	2 AL163203.2	NT	Homo saplens chromosome 21 segment HS21C003
4124	14024		12.36	3.2E-02	2 X94768.1	NT	H.saplens RP3 gene (XLRP gene 3)
					!	!	
4658	╛			3.25-0	2	Z	Saxiraga nidinca maturase (matv.) gene, chickopiast gene encoding chickopiast protein, partial cos
5399	- 1		1.46	3.2E-02		NT	S.griseocameum whiG-Siv gene
5399					3.2E-02 X68709.1	TN	S.griseocameum whiG-Stv gene
5931	15836	26959	2.26	3.2E-0		L	Rat/polyomavirus left junction in cell line W98.14
							yd33h12.s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:110087 3' similar to contains
2882	_[1		ļ	1	NAMUL -	Au repainve agnient, van inspannye eremen ,
2989	- [26013		3.25-02	AF1/3845.1	-2	Seguinus oedipus tissue kallikrein gene, complete cas
6830	16709		3.48	3.2E-02	3.2E-02 6680565 NT	5	Mus musculus kinesin family member 3c (Kif3c), mRNA
-			6				2g54b12.s1 3oares_pineal_gland_N3HPG Homo sapiens cDNA clone IMAGE:397151 3' similar to
88	١		3.82			HOMAN	BOLLOGA44 CT LOCHKUME C OXIDASE POLITIEF !! (HOMAIN);
8952	Į		1.38	3.2E-02	AB021684.1	Į	Marchantia polymorphia genes for 26S rRNA, 5S rRNA, 18S rRNA, 5.8S rRNA and 26S rRNA
1239			2.05	3.1E-02	4503418	Ę	Homo sapiens dual specificity phosphatase 4 (DUSP4) mRNA
1283			1.26	3.1E-02 P18845		SWISSPROT	NEURONAL ACETYLCHOLINE RECEPTOR PROTEIN, ALPHA-3 CHAIN PRECURSOR (GF-ALPHA-3)
1850		21621	1.36	3.1E-02	3.1E-02 6671664 NT	۲۷	Mus musculus adaptor-related protein complex AP-3, delta subunit (Ap3d), mRNA
1831	11828		1.09	3.1E-02		<u>k</u>	Drosophila melanogaster mRNA for headcase protein
	İ						

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Table 4
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Probe SEG ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Tcp Hit Descriptor
2882	12889	22688	1.07		4.1	EST_HUMAN	601338428F1 NIH_MGC_63 Homo saplens cDNA clone IMAGE:3680695 5'
3851	13782	23555	0.92	2.9E-02	H72805.1	EST HUMAN	yu07e10.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clons IMAGE:233130 5
4950		24593	1.32	2.9E-02	X65137.1	NT	S.vulgare pepC gene for PEP carboxylase
4950			1.32	2.9E-02	X65137.1	ΙN	S.vulgare pepC gene for PEP carboxylase
5799			6.47	2.9E-02	BF032233.1	EST_HUMAN	601462661F1 NIH_MGC_68 Homo sapiens cDNA clone IMAGE:3856598 5'
6298	16162	26319	10.33	2.9E-02		EST_HUMAN	601140729F1 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:3049830 5'
7568	17419	27635	1.94	2.9E-02	AW875979.1	EST_HUMAN	CM3-PT0014-071299-051-c04 PT0014 Homo sapiens cDNA
7588	17419	96922	1.94	2.9E-02	AW875979.1	EST_HUMAN	CM3-PT0014-071299-051-c04 PT0014 Homo septens cDNA
553	10494		0.87	2.8E-02	AW970153.1	EST_HUMAN	EST382234 MAGE resequences, MAGK Homolsaplens cDNA
3321	13241	23046	1.27	2.8E-02	AF066063.1	LΝ	Homo sapiens retinal fascin (FSCN2) gene, exon 2
3321	13241	23047	1.27	2.8E-02	AF066063.1	NT	Homo saplens retinal fascin (FSCN2) gene, exon 2
4214	14112		0.98	2.8E-02	8393751 NT	ĽΝ	Rattus norvegicus microtubule-associated protein tau (Mapt), mRNA
5373	15293	26140	11.41	2.8E-02	BE741083.1	EST_HUMAN	601594078F1 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:3948067 5'
6847	16726		1.33	2.8E-02	AJ005820.1	LN	Craterostigma plantagineum mRNA for homeodomain leucine zipper protein (hb-1)
9674	19574		1.51	2.8E-02	R06966.1	EST_HUMAN	y/12h02.r1 Soares fetal liver spleen 1NFLS Hamo sapiens cDNA clone IMAGE:126875 5'
0896	19237		1.33	2.8E-02	X06322.1	⊥N	Yeast CN31C chromosome III RAHS DNA (right arm transcription hot-spot)
							Human germline T-cell receptor beta chain Dopamine-beta-hydroxylase-like, TRY1, TRY2, TRY3,
							TCRBV27S1P, TCRBV22S1A2N1T, TCRBV9S1A1T, TCRBV7S1A1N2T, TCRBV5S1A1T, TCRBV13S3,
				_			TCRBV6S7P, TCRBV7S3A2T, TCRBV13S2A1T, TCRBV8S2A2PT, TCRBV7S2A1N4T,
1473			1.26	2.7E-02	U66059.1	L	TCRBV13S9/13S>
3385		23103	2		.2	TN	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 6
4107	14007	23783	2.07	2.7E-02	N47268.1	EST_HUMAN	yy86h12.r1 Soares_multiple_sclerosis_2NbHMSP Homo sapiens cDNA clone IMAGE:280487 5'
4107	14007	23784	2.07	2.7E-02	N47258.1		yy86h12.r1 Scares_multiple_sclerosis_2NbHMSP Homo saplens cDNA clone IMAGE:280487 5
6205	15985		1.9	2.7E-02		EST_HUMAN	ot96h03.s1 Scares_total_fetus_Nb2HF8_9w Homo saplens cDNA clone IMAGE:1624661 3'
9848			1.62	2.7E-02	BF514858.1	EST_HUMAN	UI-H-BW 1-anj-f-05-0-UI.s1 NCI_CGAP_Sub7 Homo sapiens cDNA clone IMAGE:3082520 3'
259	10499	20305	1.14	2.6E-02		NT	Homo sapiens chromosome 21 segment HS21C082
1345	11251		1.04	2.6E-02		EST_HUMAN	IL3-CT0219-280100-062-C09 CT0219 Homo saplens cDNA
2315			2.9	2.6E-02	AA490021.1	EST_HUMAN	ab02b02.s1 Stratagene fetal retina 937202 Homo sapiens cDNA clone IMAGE:839595 3'
2317		96022	2.86	2.6E-02	6764241 NT	LN	Mus musculus histidine rich calcium binding protein (Hrc), mRNA
2317	12198		2.86	2.6E-02	6754241 NT	TN	Mus musculus histidine rich calcium binding prolein (Hrc), mRNA
2000	ĺ		ti ti	_		<u> </u>	Mus musculus MHC class III region RD gene, partial cds; Bf, C2, G94, NG22, G9, HSP70, HSP70, HSC70t, nd can both control of the control of th
88	┙		1.93	2.05-02		120	
3883			1.13	2.6E-02		EST_HUMAN	xj68t09.x1 Soares_NFL_I_GBC_S1 Homo sapicns cDNA clone IMAGE:2862409 3
4517	14410		1.13	2.6E-02	2 BE968922.1	EST_HUMAN	601649877R1 NIH_MGC_74 Homo saplens cDŅA clone IMAGE:3933786 3'

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		_	_	,	-	_	-		_	_	_		_	_	,		_	_	_	_	_	_		_	_	_	_		-		_		_	_
	Top Hit Descriptor	Chicken dorsalin-1 mRNA, complete cds	Deinococcus radiodurans R1 section 151 of 229 of the complete chromosome 1	xa52b04.x1 NCI_CGAP_Sar4 Homo sepiens cDNA clone IMAGE:2570383 3' similar to SW:Y069_HUMAN	qg27f1.x1 NCI CGAP Kid3 Homo sapiens cDvA clone IMAGE:1762317 3'	601493473T1 NIH MGC_70 Hamo saplens cDivA clone IMAGE:3893578 3'	Rattus norvegicus Nerve growth factor receptor, fast (Ngfr), mRNA	Homo sapiens KIAA1070 protein (KIAA1070), mRNA	Homo sapiens chromosome 21 segment HS21C/103	2s84c02.r1 NCI_CGAP_GCB1 Homo saplens cDNA clone IMAGE:704162 5	UI-HF-BNO-akj-Φ-10-0-UI.rt NIH_MGC_50 Homo sapiens cDNA clone IMAGE:3077468 5'	602015501F1 NCI_CGAP_Bm64 Homo saplens cDNA clone IMAGE:4150944 5'	on26f06.y5 NCL_CGAP_Lu5 Homo saplens cDNA clone IMAGE:1557827 5'	on28f08.y5 NCI_CGAP_Lu5 Homo saplens cDNA clone IMAGE:1657827 6'	601680305R2 NIH_MGC_83 Homo sapiens cDNA clone IMAGE:3950665 3'	601680305R2 NIH_MGC_83 Homo sapiens cDNA clone IMAGE:3950665 3'	Rattus norvegicus rabphilin-34 mRNA, complete odc	H.carterae mRNA for fucoxanthin chlorophyll a/c blnding protein, Fcp1	H.carterae mRNA for fucoxanthin chlorophyll a/c blnding protein, Fcp1	PM2-NN0128-080700-001-a12 NN0128 Homo sapiens cDNA	PM2-NN0128-080700-001-a12 NN0128 Homo sapiens cDNA	hi36h08 x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2934015 3'	7630609.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3284008 3' similar to contains L1.tt L1	repetitive element;	601579393F1 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:3928054 5'	602070562F1 NCL_CGAP_Bm64 Homo sapiens cDNA clone IMAGE:4213406 5	602070562F1 NCI_CGAP_Brn64 Hamo saplens cDNA clane IMAGE:4213406 5'	HYPOTHETICAL 46.7 KD PROTEIN C19G10.05 IN CHROMOSOME I	HYPOTHETICAL 46.7 KD PROTEIN C19G10.05 IN CHROMOSOME I	Bos taurus partial stat5B gene, exons 17-19	Mus musculus major histocompatibility locus class II region; major histocompatibility protein class II alpha	chain (IAalpha) and major histocompatibility protein class II beta chain (IEbeta) genes, complete cds; butyrophilin-like (NG9), butyrophilin-li>	Home sariens gene for LECT2 complete ads	וומות מלונות וויים וויים ביים וויים ביים וויים ביים ב
i	Database Source	Z	Z	EQT LINAAN	EST HUMAN	EST HUMAN	N.	Z	Z	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	Z	TN	Z	EST_HUMAN	EST_HUMAN	EST_HUMAN		EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	SWISSPROT	SWISSPROT	Z		뉟	LZ	
	Top Hit Acession No.	L12032.1	AE002014.1	AW24454 4	AI206030.1	BE621748.1	6981271 NT	11432020 NT	AL163303.2	AA279351.1	AW500547.1	BF343827.1	AI793130.1	AI793130.1	BE974314.1	BE974314.1	U12571.1	X99697.1	X99697.1	BE701165.1	BE701165.1	AW592114.1		BE670128.1	BE746888.1	BF528722.1	BF526722.1	Q10335	Q10335	AJ237936.1		AF050157.1	ABOUTEAR 4	יייביי ייים
Most Similar	(Top) Hit BLAST E Value	2.6E-02	2.6E-02	2 80			2.6E-02	2.6E-02	2.6E-02	2.6E-02	2.6E-02	2.6E-02	2.5E-02	2.6E-02	2.5E-02		2.6€-02	2.5E-02	2.5E-02	2.6E-02	2.5E-02	2.5E-02			2.5€-02	2.5E-02	2.5€-02	2.6E-02	2.5E-02	2.5E-02		2.5E-02	0 4E 02	17.7
	Expression Signal	3.69	1.87	200	6.85	2.08	60.9	1.19	4.87	2.1	1.89	1.55	1.45	1.45	12.83	4.77	2.59	4.4	4.4	1.09	1.09	5.07		4.48	3.86	1.57	1.57	245	2.45	3.38		3.33	1 74	
	ORF SEQ ID NO:	24481	24637	7487		25895		27512	28038		28949		20274			20828		22852	22653	23643	23644	23788				26555	26556	28320		28370				
Exon	SEQ ID	14695	14873	4 4809	1		15977	17306	17798	18579	18660		10463	10463	10721	10778	12597	12853	12853	15070	15070	14011		15667	15674	16378	16378	18071		18118		18135	1	ı
Probe	SEQ ID	4811	4998	5025	5771	5870	6130	7388	7948	8692	8848	9320	521	521	792	851	2735	2926	2926	3959	3959	4111		5759	5767	6219	6219	8185	8185	8238		8255	8708	3

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Tcp Hit Descriptor	Homo saplens similar to ALEX3 protein (H. sapiens) (LOC63634), mRNA	Homo sapiens mitogen-activated protein kinase idnase kinase 13 (MAP3K13), mRNA			П		DT H-2 CLASS I HISTOCOMPATIBILITY ANTIGEN, K-B ALPHA CHAIN PRECURSOR (H-2K(B))	T.thermophila calcium-binding 25 kDa (TCBP 25) protein mRNA, complete cds				za35g11.s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:294596 3' similar to Ab (K02909)RATSR7K Rat (RNA);contains A3R,b1 A3R repetitive element;	П	nh07b12.s1 NCI_CGAP_Thy1 Homo saplens cDNA clone IMAGE:943583 similar to contains Alu repetitive element; contains element PTR5 repetitive element;	Mus musculus major histocompatibility locus class III regions Hsc70t gene, partial cds; smRNP, G7A, NG23, Muss homdog, CLCP, NG24, NG25, and NG26 genes, complete cds; and unknown genes		Mus musculus major histocompatibility locus class III regions Hsc70t gene, partial cds; smRNP, G7A, NG23, Muts homolog, CLCP, NG24, NG25, and NG26 genes, complete cds; and unknown genes	Bacterlophage bIL67, complete genome	Mus musculus DinB homolog 1 (E. coli) (Dinb1), mRNA	П	П	Caenorhabdilis elegans mRNA for Iron-sulfur subunit of mitochondrial succinate dehydrogenase, complete cots	Т	Т	Т	П	\neg	4 Homo sapiens mammary tumor-associated protein INT6 (INT6) gene, exon 4	S.cerevisiae chromosome IV reading frame ORF YDL 245c
Top Hit Database Source	TN	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	SWISSPROT	SWISSPROT	TN	SWISSPROT	SWISSPROT	LN.	EST_HUMAN	EST HUMAN	EST_HUMAN	<u> </u>		LN LN	TN	NT	EST_HUMAN	NT	ĻΝ	EST HIMAN	PACE FOLLOWING	NEW CONTRIB	SWISSPROT	EST_HUMAN	LN	Ŀ
Top Hit Acesslan No.	11420078 NT	11433220 NT	BE973327.1	AI378582.1	H65884.1	P01901	P01901	J05110.1	P01901	P01901	AL161595.2	N69442.1	AV692954.1	AA493894.1	AF109905.1		AF109905.1	P627909 NT	6753635 NT	BE928869.1	AF163864.1	ARANGERS 1	MA2080 4	N#2300.1	Bro/84//.	P54643	W05340.1	U94165.1	274293.1
Most Similar (Top) Hit BLAST E Value	2.5E-02	2.5E-02	2.6E-02	2.4E-02	2.4E-02	2.4E-02	2.4E-02	2.4E-02	2.4E-02	2.4E-02	2.4E-02	2.4E-02	2.4E-02	2.4E-02	2.4E-02		2.4E-02	2.4E-02	2.4E-02	2.4E-02	2.4E-02	2 4F-02	2 4 50	2.4E-02	2.4E-02	2.4E-02	2.3E-02	2:3E-02	2.3E-02
Expression Signal	219	1.63	2.39	1.01	1.89	2.22	2.22	4.1	1.43	1.43	11.51	10.36	217	2.98	1 88		1.96	1.95	2.72	2.03	1.27	3.87	9,	0 3	1.38	1.59	5.79	7.89	2.52
ORF SEQ ID NO:			25271	19958	21344		21785	23948	24099			27032		27808			28967		25357										22081
Exan SEQ ID NO:	19657	19528	19167	10139	11484	ſ	12710	14171	14314	ı	14987	16840	ì	<u> </u>	1	ı	18669	18856	18947	1	19036	20.20	1	-1	_l	19667	11726	11740	12183
Probe SEQ ID NO:	9280	9463	9581	167	1580	1998	1998	4272	4420	4420	5119	6962	7643	7734	RAEZ		8857	8080	9224	9282	9368	080	3	7229	9538	9977	1829	1844	2302

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Table 4
Single Exon Probes Expressed in Heart

	Top Hit Descriptor	ue Homo sapiens cDNA	cats cats	lete cds	apiens cDNA	aplens cDNA	apiens cDNA	A clone IMAGE:27706713'	'A clone IMAGE:2770671 3'	apiens cDNA	IA clone IMAGE:3955386 5	IA clone IMAGE:3955386 5'	JNA clane IMAGE:2028868 3'	ONA clone IMAGE:2028868 3'	Caulabacter crescentus topoisomerase IV ParE subunit (parE) gene, complete cds, and proplonyl-CoA	tial cds	fragment No. 17	omplete cds	of the complete gename	of the complete genome	GLÜCOAMYLASE S1/S2 PRECURSOR (GLÜCAN 1,4-ALPHA-GLÜCOSIDASE) (1,4-ALPHA-D-GLÜCAN GLÜCOHYDROLASE)	4A clone IMAGE:3546567 5'	or gene, complete cds	Dictyostelium discoideum extracellular signal-regulated protein kinase (ERK1) mRNA, complete cds	RIL1), mRNA	Columba livia nucleoside diphosphate kinase (NDPK) gene, nuclear gene encoding mitochondrial protein, complete ods	ling protein 2 (CHD2) mRNA	168	ons 1 through 4	DNA clone IMAGE:1084782 3'	II A VD1 gene complete cds
	Top Hit Database Source	EST_HUMAN HSAAACADH P, Human foetal Brain Whole tissue Homo sapiens cDNA	Gallus gallus connexin 45.8 (Cx45.8) gene, complete cds		EST_HUMAN CM4-NN0080-290400-160-b04 NN0080 Homo sapiens cDNA	EST_HUMAN CM3-MT0118-010900-318-g07 MT0118 Homo saplens cDNA	EST_HUMAN CM3-MT0118-010900-318-907 MT0118 Homo sapiens cDNA	EST_HUMAN xs25d08.x1 NCI_CGAP_U2 Homo sepiens cDNA clone IMAGE:2770671 3	EST_HUMAN xs25d08.x1 NCL_CGAP_Ut2 Homo saplens cDNA clone IMAGE.2770671 3'		EST_HUMAN 601672279F1 NIH_MGC_20 Homo sapiens cDNA clone IMAGE:3955386 5'		EST_HUMAN qz35c03.x5 NC _CGAP_Kid11 Homo sapiens c0NA clone IMAGE:2028668 3'	EST_HUMAN qz35c03.x5 NCI_CGAP_Kid11 Hc	Caulobacter crescentus topoisome						SWISSPROT GLUCOHYDROLASE)	Τ	T Streptomyces sp. alpha-1,3/4-fucosidase precursor gene, complete cds		Homo sapiens dead ringer (Drosophila)-like 1 (DRIL1), mRNA					T_HUMAN	T Infectious bursal disease virus segment B strain IL4 VP1 gene, complete cds
3.6	Top Hit Acession No.	Z20377.1 E	L24799.1	L24799.1	AW899107.1 ES	BE935225.1 ES	BE93525.1 EX			BE143150.1 ES	BF026487.1 ES	BF026487.1 ES	AI793177.1 ES	AI793177.1 ES		U86303.1 NT	AL161505.2 NT	U63610.1 NT	AE000199.1 NT	AE000199.1 NT	15 07980d	31.1	U39394.1	U11077.1	11426388 NT	AF018267.1 NT	57448	Z82001.1 NT	AF109633.1 NT	AA577785.1	AF083094 1
	Most Similar (Tap) Hit BLAST E Vatue					2.3E-02	2.3E-02	2.3E-02	2.3E-02	2.3E-02	2.3E-02	2.3E-02	2.3E-02	2.3E-02		2.3E-02	2.3E-02		2.3E-02	2.3E-02	235-02	2.3E-02	2.3E-02	2.3E-02	2.3E-02		2.2E-02	2.2E-02	2.2E-02	2.2E-02	2 2F-02
	Expression Signal	4.79	1.19	1.19	1.28	0.81	0.81	0.94	0.94	1.75	2.39	2.39	F	1		3.67	4	5.67	1.51	1.51	2.15	4.41	1.94	2.27	1.27	3.09	1.53	1.78	1.4	1.93	3.27
	ORF SEQ ID NO:	23329	23735			L	24028	24029		24155	24171	24172	24591	24592			26018			27842	28302		25219			20482	L	21752			
	Exan SEQ ID NO:	13542	13959	L	L	L	L	L	15072	14366	14384	14384	14825	14825	ı	16223	15895	16478	17614	17614	18050	1		19756	<u> </u>		L	L	12730	13306	13509
	Probe SEQ ID NO:	3628	4057	4057	4323	4347	4347	4348	4348	4472	4480	4480	4948	4948		5302	2980	8658	7764	7764	8182	9201	9713	9767	9961	720	1713	1968	2895	3388	3595

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	_	_				_		_		_				_			_		_	_			÷	_			_	- -			_
Top Hit Descriptor	S.cerevisiae chromosome IV reading frame ORI; YDL245c	AV699721 GKB Homo sapiens cDNA clone GKBAND03 3'	Arabidopsis thaliana DNA chromosoma 4, contig fragment No. 27	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 27	Homo sapiens DNA, DLEC1 to ORCTL4 gene ragion, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes,	complete cos)	Homo sapiens DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes, complete cds)	ne47h07,s1 NCI_CGAP_Co3 Homo sapiens cDNA clone IMAGE:900541 3' similar to contains Alu repetitive	element;	AV761502 MDS Hamo saplens cDNA clone MDSADG01 5'	Dictyostelium discoideum histidine kinase C (dhkC) mRNA, complete cds	Bacillus subtills cotKLM cluster, CotK (cotK), CctL (cotL), and spore coat protein CotM (cotM) genes,	complete cds	Mus musculus macrophage migration inhibitory factor (MIF) gene, 5' flanking region and partial cds	Mus musculus macrophage migration inhibitory factor (MIF) gene, 5' flanking region and partial cds	Tegula aurectincta major acrosomal protein precursor (TMAP) mRNA, complete cds	ух43h07.r1 Scares melanocyte 2NbHM Homo stiptens cDNA clone IMAGE:264541 5	PM2-BT0546-120100-001-f11 BT0546 Homo sapiens cDNA	PM2-BT0546-120100-001-f11 BT0546 Homo saplens cDNA	zx63b09,r1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA clone IMAGE:796121 5	S.cerevisiae chromosome IV reading frame ORF, YDL245c	Borrella burgdorferi plasmid cp32-2, erpC and erpD genes, complete cds; and unknown genes	wg81d11.x1 Soares_NSF_F8_9W_OT_PA_P_S1 Homo sapiens cDNA clone IMAGE:2371509 3	A.thaliana mitochondrial genome, part A	wh54e05.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2384528 3'	Homo sapiens partial 5-HT4 receptor gene, exons 2 to 5	Homo sapiens partial 5-HT4 receptor gene, exons 2 to 5	Homo sapiens putative psihHbA pseudogene for hair keratin, exons 2 to 7	Azospirillum brasilense major outer membrane protein OmaA precursor (omaA) gene, complete cds	7g61c08.x1 NCI_CGAP_Pr28 Homo saplens cENA clone IMAGE:3309998 3' similar to contains MER1.t3 MER1 repetitive element ;	QV4-NN0038-270400-187-h05 NN0038 Homo sapiens cDNA
Top Hit Database Source	NT	EST_HUMAN	NT	NT		Z	FN		EST_HUMAN	EST_HUMAN	N.		NT	NT	NT	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	NT	EST HUMAN	NT	EST_HUMAN		TN	NT	LN LN	EST_HUMAN	EST_HUMAN
Top Hit Acesslan No.	274293.1	AV699721.1	AL161515.2	AL161515.2		AB026898.1	AB026898.1		AA503553.1	AV761502.1	AF029726.1		U72073.1	AF204395.1	AF204395.1	AF190899.1	N29266.1	BE072546.1	BE072546.1	AA461271.1	274293.1	U44914.1	AI768127.1	Y08501.1	A1823432.1	AJ243213.1	AJ243213.1	Y19213.1	AF183913.1	BF002932.1	AW895565.1
Most Similar (Top) Hit BLAST E Value	2.2E-02	2.2E-02	2.2E-02	2.2至-02		2.2€-02	2.2E-02		2.2E-02	2.1E-02	2.1E-02		2.1E-02	2.1E-02	2.1E-02	2.1E-02	2.1E-02	2.1E-02	2.1E-02	2.1E-02	2.1E-02	2.1E-02	2.1E-02	2.1E-02	2.1E-02	2.1E-02	2.1E-02	2.1E-02	2.1E-02	2.0E-02	2.0E-02
Expression Signal	0.99	3.78	2.28	2.28		2.33	233		2.24	5.58	7.89		7.43	1.91	1.91	0.95	3.98	0.85	0.85	1.23	0.89	1.61	1.03	4.65	0.86	1.8	1.8	4.99	3.33	1.35	7.86
ORF SEQ ID NO:	23549	26318				27651	27652	L					20898	21125	21128	21694	20530		Ŀ	23247	23719	24022	24033	24288					26184		
Exan SEQ ID NO:	13755	16161	L	١.	1	17436	17436	L	19096	1_	L		11149	11270	11270	11815	10693	L	l _	_	13941	14239	14247	14499	14600	_	L	L_		<u> </u>	1
Probe SEQ ID NO:	3844	6297	6876	6876		7585	7685		9466	413	144		1242	1365	1365	1920	2785	3110	3110	3535	4038	4342	4361	4811	4714	7553	7553	9449	9857	17	18

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	П	\neg	٦	_	T	٦	٦	\neg	┒	7			T	٦	T	Т	╗	٦	7	\neg	T	7	T	\neg	Ţ	٦			\neg	丁	7	T	\neg
řop Hit Descriptor	Mus musculus DinB homolog 1 (E. coli) (Dinb1), mRNA	aa15b10,r1 Soares_NhHMPu_S1 Homo sapiens cDNA clone IMAGE:813307 5'	Mus musculus DinB homolog 1 (E. coll) (Dinb;), mRNA	Homo sapiens genomic region containing hypervariable minisatellites chromosome 1[1p36.33] of Homo	saplens	Homo sapiens hypothetical protein FLJ10379 (FLJ10379), mRNA	Homo sepiens hypothetical protein FLJ10379 (FLJ10379), mRNA	Homo sapiens hypothetical protein FLJ10486 (FLJ10486), mRNA	Homo sapiens hypothetical protein FLJ10486 (FLJ10486), mRNA	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 32	7g51c08.x1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:3309998 3' similar to contains MER1.t3 MER1 repetitive element ;	Mus musculus sema domain, transmembrane domain (TM), and cytoplasmic domain, (semaphorin) 6B	(Jemado), MKINA	Arabidopsis ihallana CZHZ zinc linger protein r.Zr. mKNA, complete cos	P. vulgaris hydroxyproline-rich glycoprotein (HRGP) mRNA, 3' end	Homo sapiens chromosome 21 segment HS210078	aa15b10.r1 Soares_NhHMPu_S1 Homo saplens cDNA clone IMAGE:813307 5	Japanese encephalitis virus envelope protein mRNA, partial cds	wa17b02.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2298315 3'	Mycobacterium tuberculosis H37Rv complete genome; segment 93/162	Equus caballus DNA for 17alpha-hydroxylase/17,20-lyase, complete cds	Homo sapiens ankyrin 3, node of Ranvier (ankyrin G) (ANK3), transcript variant 1, mRNA	Homo sapiens ankyrin 3, node of Ranvier (ankyrin G) (ANK3), transcript variant 1, mRNA	aa15b10.r1 Soares_NhHMPu_S1 Homo sapiens cDNA clone IMAGE:813307 5	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 32	601478819F1 NIH_MGC_68 Home saplens cDNA clone IMAGE:3881477 5'	yd04c08.r1 Soares infant brain 1NIB Homo saplens cDNA clone IMAGE:24675 51	nf19e07.s1 NCI_CGAP_Pr1 Homo sapiens cDNA clone IMAGE:914196 similar to contains L1.t1 L1	repetitive element;	EMPTY SPIRACLES HOMEOTIC PROTEIN	Homo sapiens chromosome 21 segment HS21C103	Homo sapiens chromosome 21 segment HS21C103	Arabidopsis thallana DNA chromosome 4, contig fregment No. 50
Top Hit Database Source		EST_HUMAN	L		NT	LN	NT	NT	LZ	NT	EST HUMAN	!	Z	NT	NT	NT	EST_HUMAN	LN	EST_HUMAN	NT	NT	TN	NT	EST_HUMAN	NT	EST_HUMAN	EST HUMAN		EST_HUMAN	SWISSPROT	NT	NT	NT
Top Hit Acesslan No.	6753635 NT	AA456538.1	6753635 NT		AL096805.1	TN 1622391	8922391 NT	8922453 NT	8922463 NT	AL161532.2	BF002832.1		7305474 NT	AF095588.1	M18095.1	AL163278.2	AA456538.1	U70408.1	A1640342.1	Z73966.1	D88184.1	10947055 NT	10947055 NT	AA456538.1	AL161532.2	BE786595.1	T80037.1		AA572764.1	P18488	AL163303.2	AL163303.2	AL161550.2
Most Similar (Top) Hit BLAST E Value	2.0E-02	2.0E-02	2.0E-02		2.0E-02	2.0E-02	2.0E-02	2.0E-02	2.0E-02	2.0E-02	2.0E-02			2.0E-02	2.0E-02	2.0E-02	2.0E-02	2.0E-02	2.0E-02	2.0E-02	2.0E-02	2.0E-02	2.0E-02	2.0E-02	2.0E-02		2.0E-02		1.9E-02	1.9E-02	1.9E-02		1.9E-02
Expression Signal	263	2.62	1.75		0.97	1.61	1.61	2.31	231	1.75	1.98		1.16	3.33	1.5	66.0	0.99	1.8	1.59	2.01	2.55	3.09	3.09	1.41	1.68	1.72	4.08		1.68	0.84	2.52	2.52	0.87
ORF SEQ.	20040				20830	20838		21600			19795				23814	24725				28175		28031	28032						20432	21364	21779		
Exon SEQ ID NO:	10225	1_	L		10987	11092	11092	<u> </u>	11727	<u> </u>	10001	<u> </u>	- 1	13114	13834	14949	14955	17529		L	18586	18739	18739	L	12630	L	L	Ł	10811	11504	L	L	12335
Probe SEQ ID NO:	259	292	781		1071	1181	1181	1830	1830	2768	3040		3105	3189	3925	5079	5085	7679	7930	8037	8878	8931	8931	9019	9481	8828	9935		878	1699	1993	1993	2458

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Probe SEO ID	SEQ ID	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E	Tap Hit Acession No.	Top Hit Database Source	.Ťop Hit Descriptor
<u>.</u>	<u></u>			Value			
2875	12802		7.08	1.9E-02	AA713856.1	EST_HUMAN	nw04f05.s1 NCI_CGAP_SS1 Hamo sepiens cDNA clone IMAGE:1238337 3'
2923	12850	22650	1.53	1.9E-02	AV648669.1	EST HUMAN	AV648669 GLC Homo saplens cDNA clone GLCBLH07 3'
3222	ı		0.82	1.9E-02	AB033611.1	L	Urotrichus talpoides mitochondrial gene for cytochrome b, complete cds
3560	l		1.25	1.9E-02	N52250.1	EST HUMAN	yz28b02.s1 Soares_multiple_sclerosis_2NbHMSP Homo sapiens cDNA clone IMAGE:284331.3*
3844	1		6.75	1.9E-02	BE738088.1	EST_HUMAN	601572682F1 NIH_MGC_57 Homo saplens clDNA clone IMAGE:3839564,5'
	1					Г	qn04c07.x1 NCI_CGAP_Lu5 Homo sapiens cIDNA clone IMAGE:1897260 3' similar to contains Alu repetitive
3654	13568	23355	0.98	1.9E-02	Al301183.1	T_HUMAN	element;
3961	13868	23646	1.14	1.9E-02	AF141940.1	NT	Mycoplasma imitans VihA1 precursor (vihA1) and VihA2 precursor (vihA2) genes, partial cds
4099	13999	23777	1.47	1.9E-02	P09081	SWISSPROT	HOMEOTIC BICOID PROTEIN (PRD-4)
4099	13999	23778	1.47	1.9E-02	P09081	SWISSPROT	HOMEOTIC BICOID PROTEIN (PRD-4)
							(146d04.X1 Soares_NSF_F8_9W_OT_PA_P_S1 Homo sapiens cDNA clone IMAGE:2144551 3' similar to
4441	14335	24125	2.51	1.9E-02	AI452989.1	EST_HUMAN	contains Alu repetitive element;
4944	12335	22230	1.9	1.9E-02	AL161550.2	LN	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 50
5356	15276	25108	1.29	1.9E-02	L47572.1	L	Meleagris gallopavo paraoxonaso-2 (PON2) mRNA, complete cds
6985	16862		1.29	1.9E-02	AL162754.2	LN	Neisseria meningitidis serogroup A strain Z2491 complete genome; segment 3/7
7377	l	27452		1.9E-02	BF316129.1	EST_HUMAN	601886130F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:4125462 5'
7777	L	27859	1.31	1.9E-02	BF695832.1	EST_HUMAN	601852383F1 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:4076253 5
9234	19518	25137	2.55	1.9E-02	AF101065.1	NT	Hirudo medicinalis intermediate filament gilarin inRNA, complete cds
							hn52c08.x1 NCI_CGAP_Co17 Homo sapiens cDNA clone IMAGE:3027274 3' similar to contains element
343	10302	20117	1.44	1.8E-02	AW771104.1	EST_HUMAN	MER29 repetitive element ;
670	10604	20421	1.14	1.8E-02	BF308122.1	EST HUMAN	601884329F1 NIH_MGC_17 Homo saplens CCNA clone IMAGE:4139983 5
1144	11057	20900	1.34	1.8E-02	X17664.1	NT	H.francisci mRNA for myelin basic protein (MBP)
1417	11323		1.23		AF243382.1	TN	Drosophila melanogaster cytoplasmic protein ericore (enc) mRNA, complete cds
2646	5 12612	22403	1.51	1.8E-02	AE004544.1	NT	Pseudomonas aeruginosa PA01, section 105 of 529 of the complete genome
3174	13099		0.89		AI805829.1	EST_HUMAN	te52a09.x1 Soares_NFL_T_GBC_S1 Homo saptens cDNA clone IMAGE:2090296 3'
3812	13724	23513	1.09		AW879122.1	EST_HUMAN	MR1-0T0011-280300-009-g04 OT0011 Homo sapiens cDNA
3812	L		1.09	1.8E-02	AW879122.1	EST HUMAN	MR1-OT0011-280300-009-g04 OT0011 Homo sapiens cDNA
3998	13903		1.15	1.8E-02	AA861446.1	EST_HUMAN	ak24h04.s1 Scares_testis_NHT Homo seplens cDNA clone IMAGE:1406935 3'
4332	L	24011		1.8E-02	AW936363.1	EST_HUMAN	QV4-DT0021-301299-071-b11 DT0021 Homo sapiens cDNA
4873	3 14763			1.8E-02	060810	SWISSPROT	HYPOTHETICAL PROTEIN DJ845024.2
4887	7 14767	24543	1.08		AI288701.1	EST_HUMAN	qm08b04.x1 NCI_CGAP_Lu5 Hamo sapiens cDNA clone IMAGE:1881007 3'
6075	5 16058		3.96	1.8E-02	P14310	SWISSPROT	HYPOTHETICAL 7.9 KD PROTEIN IN FIXW 5/REGION
7580	17411		2 49		AA897543.1	EST HUMAN	aj62f09.s1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1394921 3' similar to gb:L11672 ZINC FINGER PROTEIN 91 (HUMAN);
	1			1			

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					D:::)		
Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEO ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
7790	17840	27873	1.51	1.8E-02	BE778274.1	EST_HUMAN	601463545F1 NIH_MGC_67 Homo sapiens cDNA clone IMAGE:3866963 5'
7871	L			1.8E-02	X96933.1	NT	L. stagnalis mRNA for myomodulin neuropeptide precursor
8735	1_	L		1.8E-02	AB002337.2	NT	Homo sapiens mRNA for KIAA0339 protein, pertial cds
8735	١.		1.78	1.8E-02	AB002337.2	NT	Homo sapiens mRNA for KIAA0339 protein, perital cds
8888	ı.		1.78	1.8E-02	AP000006.1	NT	Pyrococcus horkoshii OT3 genomic DNA, 1165001-1485000 nr. posiuon (o//)
8897				1.8E-02	U62749.1	L	Zea mays addic ribosomal protein P2a-3 (rpp2a-3) mRNA, partial cds
9912	1.			1.8E-02	AF047475.1	LN	Drosophila melanogaster projectin (projectin) gime, partial cds
888	L	20662		1.7E-02	BE394869.1	EST_HUMAN	601310626F1 NIH MGC 44 Homo sapiens cUNA clohe IMAGE 303219U 3
	L			·			ht34s03.x1 Soares_NFL_T_GBC_S1 Home sapiens cUNA clone IMAGE:2933/40 3 similar to contains
1750	11650	21519	2.24	1.7E-02	AW573183.1	EST_HUMAN	L1.ft L1 repetitive dement;
	l _			00 24 7	A)A(679409 4	NAMILIA FOR	ht34e03.x1 Soares_NFL_1_GBC_S1 Home saptens conveniented invage.zeco.tyclolominal to conveniented to the profit of
1750		02315.			AV 373163.1		Homo sapiens chromosome 21 segment HS210004
1828			3.08			1	Describering mRNA for milistratumin 29, complete cds
2068	11856	3	13.03		AB00481	Z	Oryconegus Confederation Transaction From Secretary Promotions (RABEXS), mRNA
2603	12471	-	1.35			LN	Homo saplens putative read of the Aviange for a figure in 1875-168682 3'
8882	12895	5 22695	0.92	1.7E-02	AI147615.1	EST_HUMAN	db22a08 XI Soares pregnant urefus Nonro septients con control in the control of t
	į.				1	MAN ILL POS	hm45e04.x1 NO_CGAP_RDF1 Homo sapiens;cDNA clone IMAGE;3015554 \$ strinks to currents MFR19 b1 MFR19 repetitive element;
3467		3	4.17		_	TO COMPLY	LINETIDANIE BICH OI VOODBOTEIN PRECIESOR
3573	13487	,	0.88	1.7E-02	P04929	SWISSPROI	TICLI IDINIE-YICOTO COLOR CONTROL CONT
02.07	13080		80.0	1 7F-02	AA669618.1	EST HUMAN	actitition is strategene ovary (#837.217) runio saprens contra ciono instrucciocos. I communication repetitivo element; contains element MER24 repetitive element;
4105	L		1 78			EST HUMAN	ye86f08.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:124647 6
ř	1_				_		hf34a03.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2933740 3' similar to contains
4427	14322	2 24109	1.24	1.7E-02	AW573183.1	EST HUMAN	L1.H L1 repetitive element;
4615	1	L	1.77	1.7E-02	V00641.1	TN	Messenger RNA for anglerfish (Lophius americanus) somatostatin II
4708	Ł.				A1015076.1	EST_HUMAN	ov51e02.s1 Soares_testis_NHT Homo saplens cDNA clone IMACE:104Ucco o
4987	1	24812		1.7E-02	AF105037.1	NT.	Murid herpesvirus 4 complete genome
	丄						wg35f09.x1 Soares_NSF_F8_9W_OT_PA_P_\$1 Homo sapiens cDNA clone IMAGE:2367113 3 similar to
5728	5 15632	25735	1.53	1.7E-02	AI769247.1	EST_HUMAN	contains Alu repetitive element;
6276	<u>L</u>		3 2.31	1.7E-02	8400716 NT	INT	Homo sapiens nebulin (NEB), mRNA
6554			1.81		AJ010770.1		Homo sapiens hyperion gene, exons 1-50
7592	2 17443	3 27659	1.48	1.7E-02		EST_HUMAN	DKFZp4340314 rt 434 (synonym: ntess) nomo sapiens court cidae on appeara
9781					2 AW 903482.1	EST HUMAN	CM4-NN1030-040400-130-106 NN 1030 Homo septens CONA
9861	Ш	Ц	1.98	1.7E-02	2 Q03211	SWISSPROT	PISTIL-SPECIFIC EXTENSIN-LINE PROTEIN/PRECONSON (T.C.E.T.)

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	Top Hit Descriptor	Mycobacterium tuberculosis H37Rv complete genome; segment 13/162	Treponema maltophilum flaB2, flaB3 and fliD genes for flagellin subunit proteins and CAP protein homologue	LIVER CARBOXYLES LERASE 22 PRECURSON (EGASTIN) (ESTEPASE 22)	LIVER CARBOXYLES I ERASE 22 PRECORSON (EGASTIV) (ESTENASE 22)	Homo saplens KVLQ in gene	ne81d06.s1 NCI_CGAP_Ew1 Homo sapiens cDNA clone invACE.3 loco/	Homo sapiens mRNA for KIAA0634 protein, partial cos	IL3-C10219-160200-063-C07 C10219 Homo tapiens CUNA	Mus musculus major histocompatibility complex region NG27, NG28, NF528, NADH oxoodecudase, NG29, KIFC1, Fas-binding protein, BING1, tapasin, RalGDS-like, KE2, BING4, beta 1,3-galactosyl transferase, and	RPS18 genes, complete cds; Sacm21 gene, partial>	QV2-PT0012-140100-030-107 PT0012 Homo seplens cDNA	Mus musculus CD5 antigen (Cd5), mRNA	Candida albicans CaGCR3 gene, complete cds	Human apoC-II gene for preproapolipoprotein C-II	Drosophila melanogaster enhancer of polycomb (E(Pc)) mRNA, complete ods	nf19g03.s1 NCI_CGAP_Pr1 Homo sapiens cDNA clone IMAGE:914260 similar to SW:1ELO_KAB11 P29294 TELOKIN. [1];	inf19a03.s1 NCI_CGAP_Prf Homo saplens oDNA clone IMAGE:914260 similar to SW:TELO_RABIT	P29294 TELOKIN. [1]:	Gigalius microsatellite DNA (LEI0260 (= 116)III=11))	Arabidopsis thallana DNA chromosome 4, contiguration 20	Arabidopsis maliana DINA chigniosonile 4, Conto inginiani 1905. 2007.	4296e10.x1 Sogres, pregnant utelus, Nontro ripolito septens contra civilis introduces pregnant utelus per prognant utelus per prognant utelus per prognant utelus per prognant utelus per per per per per per per per per per	LIVER CARBOXYLES I ERASE ZZ PRECORSON (EGAS IN) (ESTENSON ZZ)	LIVER CARBOXYLESTERASE 22 PRECURSON (EGASTN) (ESTENAGE-22)	R.narvegicus gene for choline acetyfiransferase, exon 1 (non coning)	Homo sapiens SEC14 (S. cerevisiae)-like 2 (SEC14L2), mKNA	Homo sapiens transcription factor (HSA130894), mKNA	W27b07.s1 Soares fetal liver spieen 1NFLS Homo saplens CUNA cione livings 245525.5	Arabidopsis thaliana DNA chromosome 4, contig tragment No. 90	Homo sepiens CACNA1F gene, exans 1 to 48	Homo capiens CACINATE gene, awais 1 to 40
-	Top Hit Database Source	TN TN		╗	ISSPROT	ı	T HUMAN		EST HUMAN IL	≥ ₹	N N	EST_HUMAN Q		NT	N	D LN	EST HIMAN	Т	T HUMAN	NT		Т	EST HUMAN 9	T	ISSPROT	F.			T_HUMAN			LN.
26	Top Hit Acession No.	\L021929.1						AB014534.1	AW850652.1		AF110520.1	AW875407.1	6671715 NT	AB015281.1	X05151.1	AF079764.1	A A 577818 1	200	AA572818.1	294828.1	AL161508.2		AI373558.1	Q64176	Q64176	X92751.1	11417969 NT	8923734 NT		AL161594.2	AJ006216.1	AJ006216.1
	Most Similar (Top) Hit BLAST E Value	1.6E-02		1.6E-02		•	1.6E-02	1.6E-02	1.6E-02		1.6E-02	_				1.6E-02	4 85 02	70.70	1.6E-02	1.6E-02	1.6E-02	1.6E-02	1.6E-02	1.6E-02	1.6E-02	1.6E-02	1.6E-02	1.5E-02	<u> </u>			1.5E-02
	Expression	1.58	1.13	1.13	1.13	0.98	1.47	1.14	3.83		2.18	49.0	1:31	2.11	4.01	271	1 4		1.47	2.17	2.52	2:52	1.91	1.35	1.35	2.3	1.42	24.9		2.38	1.44	1.44
	ORF SEQ ID NO:		24399	21890	L	22284			23202			23885		L	L		<u> </u>	Z00444	28045		28683	3 28684		21990			100		21879			1 22744
	Exan SEQ ID NO:	10443	11539	12089	12089	<u> </u>	Ł	L	L		43086	L	L	L	L	L	1	1/804	17804	1	L	18416	18615	L	L	1_		1	L	ı	1	12951
	Probe SEQ ID NO:	501	1635	2202	2202	2518	2606	2662	3481		7007	4203	5458	6003	A758	7777	2	\$ \$ \$	7954	8280	8544	8544	9801	9211	9211	9577	9973	734	2085	2128	3023	3023

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Probe SEQ ID NO: NO: NO: 1683 4049 4387 6596 6600 7128 8508 8508 8608 8429 8508 8608	σ	ORF SEQ ID NO: 23365 23727 24048 25819 25819 25864 27717 27717 27717 27718	Signal Signal Signal 1.23 1.23 1.53 1.53 1.25 1.25 1.25 1.25 1.25 1.25 1.25 1.25			Top Hit Database Source Source Source EST HUMAN NT NT NT NT EST HUMAN NT EST HUMAN NT EST HUMAN EST HUMAN EST HUMAN EST HUMAN EST HUMAN NT NT NT NT NT NT NT NT NT NT NT NT NT	Top Hit Database Source EST HUMAN MR4-TN0115-080900-201-b12 TN0115 Home sapiens cDNA EST HUMAN MR4-TN0115-080900-201-b12 TN0115 Home sapiens cDNA EST HUMAN Zq40g10.71 Statagene hNT neuron (#897233) Home capiens cDNA clone IMAGE:632226 6' NT Home sapiens TESTIN 2 and TESTIN 3 genes; complete cds, alternatively spliced SWISSPROT HYPOTHETICAL CALCIUM-BINDING PROTEIN C18811.04 in CHROMOSOME I SWISSPROT HYPOTHETICAL CALCIUM-BINDING PROTEIN C18811.04 in CHROMOSOME I SWISSPROT HYPOTHETICAL CALCIUM-BINDING PROTEIN C18811.04 in CHROMOSOME I SWISSPROT HYPOTHETICAL CALCIUM-BINDING PROTEIN C18811.04 in CHROMOSOME I SWISSPROT HYPOTHETICAL CALCIUM-BINDING PROTEIN C18811.04 in CHROMOSOME I SWISSPROT HYPOTHETICAL CALCIUM-BINDING PROTEIN C18811.04 in CHROMOSOME I NT Home sapiens KIA4109 protein (KIA41099), mRNA EST_HUMAN M54510.01 Soares placenta Nb2HP Home sapiens cDNA clone IMAGE:135631 6' EST_HUMAN M54510.01 Soares placenta Nb2HP Home sapiens cDNA clone IMAGE:135631 5' EST_HUMAN M54510.01 Soares placenta Nb2HP Home sapiens cDNA clone IMAGE:135631 6' EST_HUMAN M54510.01 Soares placenta Nb2HP Home sapiens cDNA clone IMAGE:135631 6' EST_HUMAN M54510.01 Soares placenta Nb2HP Home sapiens cDNA clone IMAGE:135631 6' EST_HUMAN M54510.01 Soares placenta Nb2HP Home sapiens cDNA clone IMAGE:135631 6' EST_HUMAN M54510.01 Soares placenta Nb2HP Home sapiens cDNA clone IMAGE:135631 6' EST_HUMAN RC4.CN0049-140100-011-17 CN0049 Home sapiens cDNA clone IMAGE:135631 6' EST_HUMAN M54510.01 Soares placenta Nb2HP Home sapiens cDNA clone IMAGE:135631 6' EST_HUMAN RC4.CN0049-140100-011-17 CN0049 Home sapiens cDNA clone IMAGE:135631 6' EST_HUMAN M54510.01 Soares placenta Nb2HP Home sapiens cDNA clone IMAGE:135631 6' Chlamydophila pneumoniae AR39, section 58 of 94 of the complete genome NT Kenophilus Influenzae Rd section 115 of 163 of the complete genome NT Heamophilus Influenzae Rd section 115 of 163 of the complete genome NT Home sapiens Neish protein (LOC51225), mRNA NT Home sapiens Neish protein (LOC51225), mRNA NT H
1501	11280	22805	0.96	1.4E-02 1.4E-02 1.4E-02	AF216854.1 AV723785.1 AF160969.2	NT EST_HUMAN NT	Homo sapiens headpin gane, complete cds AV723785 HTB Homo sapiens cDNA clone HTBAHH11 5' Bifidobacterium longum Na+/H+ antiporter (nhaB), cytosine deaminase, and alpha-galactosidase (aglL) genes, complete cds; and N-acetyfglucosamine/xytose repressor protein (nagG/xy/R) gene, partial cds
3352 3439	13272 13356 13358	23074	5.21	1.4E-02 1.4E-02	AW074212.1 AL161586.2 AL161586.2	EST_HUMAN NT	xb09d09.x1 NCI_CGAP_GU1 Homo saplens cDNA clone IMAGE:2575793.3* Arabidopsis thallana DNA chromosome 4, contig fragment No. 82 Arabidopsis thallana DNA chromosome 4, contig fragment No. 82
3476 3606 4387 4387 4775			0.08 0.08 6.16 7.29 7.29 6.7		36918	NT NT EST_HUMAN EST_HUMAN EST_HUMAN	Homo sepiens coagulation factor XII (Hageman factor) (F12), mRNA Mus musculus histocompatibility 2, complement component factor B (H2-Bf), mRNA EST374761 MAGE resequences, MAGG Homo sepiens cDNA 601567403F1 NIH_MGC_21 Homo sepiens cDNA
4775	14659	24448	6.7	1.4E-02	BE733142.1	EST_HUMAN	601567403F1 NIH_MGC_21 Homo saplens cDNA clone IMAGE;3842280 5'

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Table 4
Single Exon Probes Expressed in Heart

Top Hit Descriptor	nI11c04.s1 NCI_CGAP_Br2 Homo sapiens cDNA clone IMAGE:1029990 3' similar to contains Alu repetitive element;	n/11c04.s1 NCI_CGAP_B/2 Homo saplens cDNA clone IMAGE:1029990 3' similar to contains Alu repetitive element:	Mycobacterium tuberculosis H37Rv complete genome; segment 88/162	601078239F1 NIH_MGC_12 Homo sapiens cDNA clone IMAGE:3464241 6	Human IFNAR gene for interferon alpha/beta receptor	Arabidopsis thaliana F21J9.2 mRNA, complete cds	Homo saplens sperm associated antigen 7 (SPAG7), mRNA	601556462F1 NIH_MGC_58 Homo sapiens cDNA clone IMAGE:3828335 5	Homo saplens chromosome 21 segment HS210001	602128475F1 NIH_MGC_56 Homo saplens cDNA clone IMAGE:4286203 5'	602129475F1 NIH_MGC_58 Homo sapiens cDNA clone IMAGE:4286203 5'	Mus musculus beta-sarcoglycan gene, completé cds	Homo saplens chromosome 21 segment HS21C001	Mus musculus chromosome X contigB; X-linked lymphocyte regulated 5 gene, Zinc finger protein 275, Zinc	finger protein 92, mmxq28orf	Mus musculus chromosome X contigB; X-linked lymphocyte regulated 5 gene, Zinc finger protein 275, Zinc	tinger protein 92, mmxq28ort	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 46	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 46	ow06g05.x1 Soares_parathyroid_tumor_NbHPAl Homo sapiens cDNA clone IMAGE:1846072.3' similar to	contains Alu repetitive element;	Homo sapiens human endogenous retrovirus W gagC3.37 G gag (gag) gene, complete cds	Mouse kidney androgen-regulated protein (KAP) gene, complete cds	xx34e03.x1 Soares_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:2815036 3'	xv34e03.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2815036 3'	Yeast ABP1 gene for actin binding protein	Bacillus subtilis complete genome (section 14 of 21): from 2599451 to 2812870	Human herpesvirus 6B, complete genome	Homo saplens V1b vasopressin receptor (VPR3) gene, complete cds	Oryza sativa replication protein A1 (Os-RPA1) mRNA, complete cds	H.sapiens DMA, DMB, HLA-Z1, IPP2, LMP2, TAP1, LMP7, TAP2, DOB, DQB2 end RING8, 9, 13 end 14 genes
Top Hit Database Source	EST_HUMAN e	EST HUMAN 6	T	EST_HUMAN 6		NT		THUMAN		HUMAN			TN TN		¥			NT	NT IN	Г	T_HUMAN	TN TN		EST HUMAN X	П	Υ	E L		LN LN	LN	T.S
Top Hit Acessian No.	AA559030.1	A A 559030 1	AL022073.1	BE544561.1	X60459.1	AF324985.1	11426968 NT	BE739263.1	AL163201.2	BF697081.1	BF697081.1	AF169288.1	AL163201.2		AL049866.2		2 AL049866.2	1.3E-02 AL161546.2	AL161546.2		1.3E-02 A1031593.1	2 AF156961.1	1.3E-02 M63707.1	1.3E-02 AW268563.1	1.3E-02 AW268563.1	1.3E-02 X51780.1	1.3E-02 Z99117.1	TN 6905586	1.3E-02 AF152238.1	1.3E-02 AF009179.1	1.2E-02 X87344.1
Most Similar (Top) Hit BLAST E Value	1.4E-02		1		1.4E-02		1.4E-02	1.3E-02	1.3E-02	1.3E-02	1.3E-02	1.3E-02	1.3E-02		1.3E-02		1.3E-02	1.3E-02	1.3E-02		1.3E-02	1.3E-02	1.3E-02	1.3E-02	1.3E-02	1.3E-02	1.3E-02	1.3E-02	1.3E-02	1.3E-02	1.2E-02
Expression Signal	4.62	4 82	1.82	207	4.79	1.65	1.89	0.86	241	1.89	1.99	1.41	٦		1.36		1.36	1.35	1.35		4.77	1.74	1.87	4.39	4.39	2.92	1.6	1.8	28.64	1.32	0.94
ORF SEQ ID NO:	25887	<u> </u>			28790				21684		22807		21684		24854			24856		_	26489	L	_	L	28485					24912	
Exan SEQ ID NO:	15768		1	1	L	۱_	19277	11718	11807	13102	13102	13789	11807	1_	15135		1	15114	15114		16332	16817	17712	18237	L	19705	19694	Ĺ.,	19486	<u> </u>	l l
Probe SEQ ID NO:	5862	5882	6732	7408	9128	9476	9763	1821	1912	3177	3177	3888	5176		5212		5212	6156	6156		6473	6639	7862	8360	8360	9102	9482	9286	9768	9949	206

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Table 4
Single Exon Probes Expressed in Heart

					,		
Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEO ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptar
352	10310	20128	1.74	1.2E-02	AA059299.1	EST_HUMAN	zf65g01.r1 Soares retina N2b4HR Homo sepiens cDNA done IMAGE:381840 5' similar to contains element L1 repetitive element ;
445	L	20210	1.48	1.25-02	P38898	SWISSPROT	HYPOTHETICAL 17.1 KD PROTEIN IN PURS S'REGION
Ş					01483500 4	TOT LIMON	qd68e12.x1 Soares_testis_NHT Homo sapiens' cDNA clone IMAGE:1734670 3' similar to contains L1.t1 L1 reportitive element
12/00/20	10003	20402	1 00	4 2E-02	AI 163242.1	LONIOL - LV	Homo saniens chromosome 21 segment HS210013
2000	Ŀ	l			A14/470950 4	EST HIMAN	137-70 vt Shares NFI T GRC St Home satisfans cDNA clone IMAGE: 2659432 3'
2383	122/1				`	EST HIMAN	601088408F1 NIH MGC 10 Hamo saplens CDNA clone IMAGE:3454608 5
2448	1.					EST HUMAN	601068406F1 NIH MGC 10 Homo saplens cDNA clone IMAGE:3454608 5'
2596	1_				_	EST HUMAN	xj37e09.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2659432 3'
3063	L	L	6.18			EST_HUMAN	zm88e03.r1 Stratagene ovarian cancer (#937219) Homo sapiens cDNA clone IMAGE:545020 5'
3250	L	22971	1.89	1.2E-02	R62805.1	EST_HUMAN	y11b08.s1 Soares placenta Nb2HP Homo sepiens cDNA clone IMAGE:138903 3'
							Human hereditary haemochromatosis region, histone 2A-like protein gene, hereditary haemochromatosis
4830	14712	24495	2.04		U91328.1	NT	(HLA-H) gene, RoRet gene, and sodium phosphate transporter (NPT3) gene, complete cds
4974	14849		1.73	1.2E-02	AB019786.1	NT	Oynops pyrrhogaster CpUbiqT mRNA, partial cds
5017	14891	24659	2.12	1.2E-02	AV731704.1	EST_HUMAN	AV731704 HTF Homo sapiens cDNA clone HTFBHG11 5'
5516	15434		1.96	1.2E-02	D78589.1	NT	Rana rugosa mRNA for calrediculin, complete cds
6180	16066	26216	5.33	L	AF175412.1	INT	Mus musculus DNA methyltransferase (Dnmt1) gene, exons 2, 3, 4, and 5
6327	16190			1.2E-02	AV732093.1	EST_HUMAN	AV732083 HTF Homo sapiens cDNA clone HTFBJC09 6'
	l				10 C	TOGGOOMO	GMP-N-ACETYLNEURAMINATE-BETA-GALACTOSAMIDE-ALPHA-2,3-SIALYLTRANSFERASE (BETA- GALACTOSIDE ALPHA-2,3-SIALYLTRANSFERASE) (ALPHA 2,3-ST) (GAL-NAC6S) (GAL-BETA-1,3- CAI NAC AI DHA 2,3-SIALY TPANSFERASE) (ST3CAI A 2) (SIATA-R)
6765	16644		122	1.2E-02		TN TN	Homo sapiens fringe protein mRNA, partial cds
6765	L	26833			AF193612.1	N	Homo sapiens fringe protein mRNA, partial cds
7152	1		1.17	1.2E-02	T76987.1	EST_HUMAN	yd72c08.s1 Soares fetal liver spleen 1NFLS Horno sapiens cDNA clone IMAGE:113774 3'
7557		27623	2.45		_	LN	Norwalk-like virus genogroup 2 gene for capsld protein, complete cds
7575	ļ	27640	1.23	1.2E-02		TN	Homo saplens Spast gene for spastin protein
9177	ı		1.78	1.2E-02	015534	SWISSPROT	PERIOD CIRCADIAN PROTEIN 1 (CIRCADIAN PACEMAKER PROTEIN RIGUI) (HPER)
9288	19575		1.5			SWISSPROT	COLLAGEN ALPHA 1(IV) CHAIN PRECURSOR
99/6	<u> </u>		3.47			EST HUMAN	C18119 Human placenta cDNA (TFujlwara) Homo sapiens cDNA clone GEN-557G08 51
1248	11155					EST_HUMAN	zm69e11.s1 Stratagene neuroepithelium (#937231) Homo sapiens cDNA clone IMAGE:530924 3'
1679	11581	21451	1.99			N⊤	H.saplens LIPA gene, exon 4
1679	11581		1.99	1.1E-02	X75491.1	N	H.seplens LIPA gene, exon 4

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_		_	\neg	_		_	_	-1	\neg	_	$\neg \tau$		┰	┰			\neg	┰	_		Т	т	т	т	т	т	Т	т	т	Т	Т	٦
טווטופ באסוו דוסטפע באט פענפט ווו חפמו נ	Top Hit Descriptor	602018037F1 NCI_CGAP_Bm67 Homo sapiens cDNA clone IMAGE:4153808 5'	za40e05.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:295040 5	Homo sapiens MHC class 1 region	tq95b10.x1 NCI_CGAP_0v23 Hamo sepiens cDNA clone IMAGE:2216539 3' similar to SW:XPF_HUMAN Q92889 DNA-REPAIR PROTEIN COMPLEMENTING XP-F CELL ;	PM3-HT0175-300999-001-h06 HT0175 Homo saplens cDNA	RC3-ST0197-120200-016-g11 ST0197 Homo saplens cDNA	DKFZp586E0924_s1 586 (synonym: hute1) Homo sapiens cDNA clone DKFZp586E0924	QV2-ST0296-150200-028-c11 ST0296 Homo saplens cDNA	RC1-HT0256-100300-016-h07 HT0256 Homo sapiens cDNA	NEUROGENIC LOCUS NOTCH 3 PROTEIN	zn24e01.r1 Stratagene neuroepithelium NT2RAi/il 937234 Homo saplens cDNA clone IMAGE:548328 5	EST186494 Colon carcinoma (HCC) cell line II Homo sapiens cDNA 5' end	Homo saplens T-box 5 (TBX5), mRNA	ab77f1.s1 Stratagene fetal retina 937202 Homo sapiens cDNA clone IMAGE:853005 3' similar to contains	Alu repetitive element	MR3-CT0176-111099-003-e10 CT0176 Homo sapiens cDNA	CM2-HT0177-041099-017-h12 HT0177 Homo sapiens cDNA	cc22h08.s1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:1350495 3	RC0-FN0025-250500-021-d02 FN0025 Homo sapiens cDNA	601649967R1 NIH_MGC_74 Homo sapiens cDiyA clane IMAGE:3933689 3'	MR0-CT0060-081099-003-h10 CT0060 Homo saplens cDNA	HA0921 Human fetal liver cDNA library Homo sapiens cDNA	Mus musculus corticotropin releasing hormone receptor 2 (Crhr2), mKNA	yq54h01.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:199633 5	Homo saplens attractin precursor (ATRN) gene, exon 25 and complete cds, alternatively spliced	601486286F1 NIH_MGC_69 Homo sapiens cDNA clone IMAGE:3888908 5	MR4-BT0356-070100-201-h01 BT0356 Homo sapiens cDNA	MR4-BT0356-070100-201-h01 BT0356 Homo sapiens cDNA	Z.mays U3snRNA pseudogene	601459570F1 NIH MGC 66 Homo sapiens cDNA clone IMAGE: 3863177 5	(8014395/0F1 NIM_MGC_bb Homo sapiens dulyA cigne invidor.35001177
gie Exon Prob	Top Hit Database Source	EST HUMAN	EST HUMAN	NT	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	SWISSPROT	EST_HUMAN	EST_HUMAN	TN		EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	N	EST HUMAN	<u>N</u>	EST_HUMAN	EST_HUMAN	EST_HUMAN	٦	EST_HUMAN	EST_HUMAN
UIO.	Top Hit Acession No	BF345263.1	N99523.1	AF055068.1	A1653508.1	BE144637.1	AW813786.1	AL048383.2	AW820281.1	BE149611.1	Q61982	AA082578.1		11435505 NT		AA668239.1	AW846120.1	AW368128.1	AA806389.1	BE835556.1		AW845621.1	A1085086.1	. 6753521 NT	R96567.1	AF218910.1	BE876539.1	1.0E-02 AW577113.1	AW677113.1	1.0E-02/Z29642.1	1.0E-02 BF036331.1	BF036331.1
	Most Similar (Top) Hit BLAST E Value	1.1E-02	1.1E-02	1.1E-02	1.1E-02	1.1E-02	1.1E-02	1.1E-02	1.1E-02	1.1E-02	1.1E-02	1.1E-02	1.1E-02	1.1E-02		1.1E-02	1.0E-02	1.0E-02	1.0E-02	1.0E-02	1.0E-02	1.0E-02	1.0E-02	1.0E-02	1.0E-02	1.0E-02	1.0E-02	1.0E-02	1.0E-02	1.0E-02	1.0E-02	1.0E-02
	Expression Signal	4.35	3.53	10.39	2.48	0.87	76.0	1.81	0.92	2.28	6.87	2.25	4.1	3.38		2.23	3.35	16.0	1.2	2.37	1.18	66.0	0.79	4.28	2.38	0.85	0.82	2.74	2.74	2.06	4.65	4.65
	ORF SEQ ID NO:	21778		22660	23200		-	24395		26506	27230	27785	27900	28476			19785	21270		22771	22950		23505	24345	24407	24633		25766		26178	27475	
	Exon SEQ ID NO:	11886	12776	12860	13384	13846	13921	14609	14704	18339	17037	17580	17660	18224		18845	3882	11411	1	12978	13150	13380		14553	1	14870	15016	15859	15659	16037	1	17270
	Probe SEQ ID NO:	1992	2848	2933	3478	3937	4016	4723	4821	6480	7160	7710	7810	8347		9084	9	1508	2525	3051	3226	3464	3805	4687	4736	4995	5149	5751	5751	6054	7403	7403

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Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Тор Hit Descriptor
8586	18454		2.1	1.0E-02.A	AF157559.1	F	Crithidia fasciculata 27 kDa guide RNA-binding protein mRNA, complete cds; mitochondrial gene for mitochondrial product
8674	L	28846		1.0E-02/	AV760016.1	EST_HUMAN	AV760016 MDS Homo septens cDNA clone MJSBDC10 5
9142	19762			1.0E-02		SWISSPROT	SPLICEOSOME ASSOCIATED PROTEIN 62 (SAP 62) (SPLICING FACTOR 3A SUBUNIT 2) (SF3A66)
9202	19535	25061	2.95	1.0E-02	1.0E-02 AW935521.1	EST_HUMAN	RC2-DT0007-120200-016-h02 DT0007 Homo sepiens cDNA
9217	Ĺ	ŀ	5.22	1.0E-02		LN	Homo sapiens renal dipeptidase (RDP) gene, complete cds
9745	1_		244	1.0E-02	1.0E-02 X62654.1	NT	H. sapiens gene for Me491/CD63 antigen
1	<u> </u>				4 17064 00 4	NAMIN FOR	wh42f09.X1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2383433 3' similar to contains element MER22 MER22 reputitive element
6/8		Legnz	7:1			FOT CLIMAN	Entatopase NIH MGC 67 Home saniens cDNA clone MAGE:3873346 5
1243	_L		/A.L	9.0E-03 I	AL 464550 2	ES - TOWNS	Azahidosis Ihaliana DNA chromosoma 4. contin fragment No. 59
2340	12220	52123		8.05-03		-	
2593	12463	22365	1.08	9.0E-03	AJ243727.1	N	Oncorhynchus nerka proviral gypsy retrotransposon pertial reverse transcriptase and protease genes (pol- gene)
1	<u></u>				4 TOTEACT A	FN	Oncorhynchus nerka proviral gypsy retrotransposon partial reverse transcriptase and protease genes (pol
2876	12803	22500			LIQ	EST HUMAN	qh90f09.xf Soares_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:1854281 3'
2078	L					EST HUMAN	ah80f09.x1 Soares NFL_T GBC_S1 Home sapiens cDNA clone IMAGE:1854281 3'
2847	<u>.L</u>					NT	S.acidocaldarius thermopsin gene, complete cd:
4914		1				EST HUMAN	1244e10.y1 NCI_CGAP_Bm52 Hamo sapiens cDNA clone IWAGE:2291466 5
5181					1~	LN	Sargocentron sp. mixed lineage leukemia-like protein (MII) gene, partial cds
2669	L		4.17		BE745988.1	EST_HUMAN	601573438F1 NIH_MGC_9 Homo saplens cDNA clone IMAGE:3834762 5'
6597			1.18	9.0E-03	AL039991.1	EST_HUMAN	DKFZp434L0412_r1 434 (synonym: htes3) Homo saplens cDNA clone DKFZp434L0412 5'
7673	17523	. 27749	1.47	9.0E-03	P20908	SWISSPROT	COLLAGEN ALPHA 1(V) CHAIN PRECURSOR
8354			1.95		Y18000.1	NT	Homo sapiens NF2 gene
9352	19763		1.34			EST_HUMAN	PM1-HT0452-291289-001-e09 HT0452 Homo capiens cDNA
9557			11.59	9.0E-03	BE348385.1	EST_HUMAN	hw/17b09.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3183161 3'
9562	L		1.27		AF137240.1	NT	Sargocentron sp. mixed lineage leukemia-like protein (Mil) gene, partial cds
9649	18215	25259		9.0E-03	AL163267.2	TN	Homo sapiens chromosome 21 segment HS21C067
9843	18341		27.64	9.0E-03	BF351141.1	EST_HUMAN	PM1-HT0452-291299-001-e09 HT0452 Homo sapiens cDNA
1	<u> </u>		0	00 20 0	4 4 700000 4	CCT LINAMI	zh30e03.s1 Soares_pineal_gland_N3HPG Homo sapiens cDNA clone IMAGE:413596 3' similar to contains Ain renetitive element
973	10897	20744	45.2			NT	Homo sapiens adenylosuccinate lyase gene, complete cds
25	1					NT	Homo septens chromosome 21 segment HS21C083
2113	- 1	ļ		١			

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Table 4
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Probe	Exan		L	Most Similar		Top Hit	
SEQ ID	<u> </u>	OKF SEC ID NO:	Signal	(10p) Hit BLAST E Value	l op Hit Acessian No.	Database	Top Hit Descriptor
2830	12857		0.86	8.0E-03	U47048.1	FZ	Eschertehla coli microcin 24 region, DNA bindirig protein (mdbA), immunity protein (mtfl), microcin 24 (mtfS), and microcin transport protein (mtfA, mtfB) genes, complete cds
3270	13191	22890	26.0		BE171225.1	EST_HUMAN	RC1+HT0545-120200-011-b09 HT0545 Homo saplens cDNA
3314	13235		0.89	8.0E-03	AJ131016.1	TN	Homo sapiens SCL gene locus
3792	13704		1.07	60-30.8	AF058764.1	NT	Xenopus laevis bone morphogenetic protein 4 (BMP-4) gene, complete cds
4288	14185	23966	5.12	8.0E-03	BF363327.1	EST_HUMAN	CM4-NN0119-300600-223-505 NN0119 Hamo sapiens cDNA
							Mus musculus major histocompatibility complex region NG27, NG28, RPS28, NADH oxidoreductase, NG29, KFC1 Fass-binding nonlein BING1 tensein RaidDS-like KF2 RING4 beta 1.3-natarchasy transferase and
5391	15310	25164	2.7	8.0E-03	AF110520.1	TN.	RPS18 genes, complete cds; Saom21 gene, perdal>
5761	19453	25775	1.35	8.0E-03	AP000002.1	N	Pyrococcus harkoshii OT3 genomic DNA, 287001-544000 nt. position (2/7)
6048	16951	26081	4.52	8.0E-03	P65577	SWISSPROT	PROBABLE PEPTIDASE Y4NA
6229	16143	26299	2.19	8.0E-03	M17197.1	N FN	A.californica (marine gastropod mollusc) neuropeptide gene (bag cell), exon 1, 5' end
8453	16314		1.86	8.0E-03	AB038267.1	N	Turslops truncatus mRNA for p40-phox, complete cds
7166	17043	27235	3.81	8.0E-03	AW808692.1	EST_HUMAN	MR1-ST0111-111199-011-h06 ST0111 Homo capiens cDNA
7117	17567		4.72	8.0E-03	BE086509.1	EST_HUMAN	QV1-BT0677-040400-131-g03 BT0677 Homo sepiens cDNA
8148	18036	28284	1.92	8.0E-03	BE788441.1	EST_HUMAN	601475619F1 NIH_MGC_68 Homo sapiens cDNA clone IMAGE:3878405 5'
8353	18230		2.78	8.0E-03	249652.1	NT	S.cerevisiae chromosome X reading frame ORF YJR162w
8887	18575	28858	1.75	8.0E-03	AA828817.1	EST_HUMAN	od80a09.s1 NCI_CGAP_Ov2 Homo sapiens cDNA clone IMAGE:1374232
8687	18575	28859	1.75	8.0E-03	AA828817.1	EST_HUMAN	od80a09.s1 NCI_CGAP_Ov2 Homo sapiens cDNA clone IMAGE:1374232
8963	18760	29053	4.83	8.0E-03	AF064589.1	NT	Homo sapiens melanoma-associated antigan (MAGE-C1) gene, complete cds
9075	18852		1.34	8.0E-03	M69035.1	NT	Oryctolagus cuniculus elF-2a kinase mRNA, complete cds
							Homo sapiens ABCG1 gene for ABC transporter (ATP-binding cassette, sub-family G (WHITE), member 1),
1718	10007	20433	3.30	8.0E-03	AB038101.1	Z	Conferential menor HC-40 ness complete of
8	L		77 58	7 OF AS	AF097183 1	F	Contosporidium paraum HC-10 gene complete cds
ક	L	L	204	7 0F-03	AF243378 1	L _Z	Glycina max clufathlona S.transferase GST 21 mRNA partial cds
1100			2.6		AV731712.1	EST HIMAN	AV731712 HTF Homo sepiens cDNA clone HTFAZF10 5
	L						FORKHEAD BOX PROTEIN D3 (HNF3/FH TRANSCRIPTION FACTOR GENESIS) (HEPATOCYTE
1341	11247		1.19	7.0E-03	Q61060	SWISSPROT	NUCLEAR FACTOR 3 FORKHEAD HOMOLOG 2) (HFH-2)
1373	11279	21135	13.73	7.0E-03	AA668298.1	EST_HUMAN	ab79b09.s1 Stratagene fetal retina 937202 Homo saplens cDNA clone IMAGE:853145 3'
1486	11391	21251	4.8	_	AW303599.1	EST_HUMAN	xv21b02.x1 Soares_NFL_T_GBC_S1 Homo saplens cDNA done IMAGE:2813739 3'
2210	l		2.24	7.0E-03	P04929	SWISSPROT	HISTIDINE-RICH GLYCOPROTEIN PRECURSOR
3702	13616	23400	1.27	7.0E-03	AW44463.1	EST HUMAN	UI-H-Bi3-akb-c-10-0-UI.s1 NCI_CGAP_Sub5 Homo sapiens cDNA okne IMAGE:2733691 3'

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ORF SEQ Expression (Top) Hit Acession ID NO: Signal Value Source	23438 1.01 7.0E-03 AF196344.1 NT Rattus norvegicus neuronal nicotinic acetylcholine receptor subunit (Alpha10) mRNA, complete ods	1.23	1.01 7.0E-03 AW 630888.1 EST_HUMAN hh89a05.y1 NCI_CGAP_CU1 Homo saplens cDNA clone IMAGE:2969936 5'		24683 1.1 7.0E-03 BE044191.1 EST_HUMAN 093434 RETICULOCALBIN.;	24684 1.1 7.0E-03 BE044191.1 EST HUMAN 093434 RETICULOCALBIN. ;	5.07 7.0E-03 AW861059.1	28831 1.49 7.0E-03 W68251.1 EST_HUMAN 2433110.r1 Soares_fetal_heart_NbHH19W Homo capiens cDNA clone IMAGE:342476 6	3.46 7.0E-03 AA327129.1 EST_HUMAN	26980 1.3 7.0E-03 BE867386.1 EST_HUMAN Q13387 HYPOTHETICAL PROTEIN 384D8_2; contains TAR1.t2 TAR1 TAR1 repetitive element;	28127 2.39 7.0E-03 BE928133.1 EST_HUMAN GM2-CT0478-230800-347-b11 CT0478 Homo gapiens cDNA	26468 4.98 7.0E-03 [235838.1 NT S.cerevisiae chromosome II reading frame ORF/YBL077w	4.98 7.0E-03 Z35838.1 NT	BE175667.1 EST_HUMAN	27670 2.63 7.0E-03[P48982 SWISSPROT BETA-GALACTOSIDASE PRECURSOR (LACTASE)	27671 2.53 7.0E-03 P49982 SWISSPROT BETA-GALACTOSIDASE PRECURSOR (LACTASE)	1.27 7.0E-03 AV887379.1 EST_HUMAN AV687379 GKC Homo sapiens cDNA clone GKCAFC07 67	28337 3.36 7.0E-03 AB008652.1 NT Bos taurus mRNA for NDP52, complete cds	y/15h01.s1 Soares fetal liver spleen 1NFLS Homo sepiens cDNA clone IMAGE:242833 3' similar to contains	7 0E-03 BE263253 1 EST HUMAN	7.0E-03 AW868110.1 EST HUMAN	6.0E-03 AW 511148.1	20980 9.29 6.0E-03 AW611148.1 EST_HUMAN SW.PXR_HUMAN O75469 ORPHAN NUCLEAR RECEPTOR PXR:	22499 1.09 6.0E-03 AF112374.1 NT Danio rerio odorant receptor gene cluster	3.11 6.0E-03 AA769135.1 EST_HUMAN	22582 3.11 6.0F-03 A 7561:35 1 FST HUMAN lah78e11.s1 Soares testis NHT Home sapiens cDNA clone 1321772.3'
	23436				24683	24684		25831	25967	26980	26127	26466	26467	26792	27670	27671		28337				20979	20980	22499	22581	22582
Exon SEQ ID NO:	13653	14329	14389		14910	14910	19452	15718	15844	16859	16992	16302	16302		17458	17456	17758	18086	10750	1	•	1	11129	12606		L
Probe SEQ ID NO:	3741	4434	4485	4916	6038	5038	5717	5813	5939	6954	6210	8441	6441	6723	7605	7605	7906	8201	Copo	8615	8962	1220	1220	2744	2862	0000

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Top Hit Descriptor	yr77h04,r1 Soares fetal liver spleen 1NFLS Hamo sapiens cDNA clone IMAGE:211351 5'	Notoncus sp. cytochrome c oxidase subunit II gene, partial ods; mitochondrial gene for mitochondrial product	Fugu rubripes zino finger protein, isotocin, fatty/acid binding protein, sepiapterin reductase and vasotocin genes, complete cds	Fugu rubripas zinc finger protein, isotocin, fatty acid binding protein, sepiapterin reductase and vasotocin genes, complete cds	2c13a11.r1 Soares_parathyroid_tumor_NbHPA/Homo sapiens cDNA clone IMAGE:322172 6'	UI-H-BI4-apm-c-08-0-UI.s1 NCI_CGAP_Sub8 Homo sapiens cDNA clone IMAGE:3087754 3'	Mus musculus glucosamine-6-phosphate deaminase (Gnpi), mRNA	RC0-CT0204-240999-021-b10 CT0204 Homo sapiens cDNA	600942904F1 NIH_MGC_15 Homo sapiens cDNA clone IMAGE:2959513 5'	ly62h10.s1 Soares_multiple_sclerosis_2NbHMSP Homo sapiens cDNA clone IMAGE:278179 3'	ov33c11.x1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1639124 3'		Homo sapiens chromosome 21 segment HS21C081	aj95509 s1 Soares_parathyroid_tumor_NbHPA Homo sapiens cDNA clone IMAGE:1404256 31	Homo sapiens telomerase reverse transcriptase (TERT) gene, exons 7-16 and complete cds	ow13a04.x1 Soares, parathyroid, tumor_NbHPA/Homo sapiens cDNA clone IMAGE:1646670.3' similar to contains MER10.b1 MER10 repetitive element;	RC0-UM0051-210300-032-g02 UM0051 Homo seplens cDNA	601454915F1 NIH_MGC_66 Homo saplens cDNA clone IMAGE:3858626 5'	Subacute sclerosing panencephalitis (SSPE) virus mRNA for fusion protein	## 02202.x1 NOI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2131202 3' similar to SW:R13A_HUMAN P40429 60S RIBOSOMAL PROTEIN L13A;	M.thermoformicicum complete plasmid pFV1 DNA	Г	Γ	te91c12:x1 NCI_CGAP_Pr28 Homo sapiens cDINA clone IMAGE:2094070 3' similar to TR:O00519 O00519 FATTY ACID AMIDE HYDROLASE.;	te91c12.x1 NCI_CGAP_Pr28 Homo saplens cDNA done IMAGE:2094070 3' similar to TR:000519 000519 FATTY ACID AMIDE HYDROLASE.	Mus musculus zinc-finger protein mRNA, complete cds	601572746F1 NIH_MGC_57 Homo sapiens cDNA clone IMAGE:3839747 5'
Top Hit Database Source	EST_HUMAN	NT	TN	Ę	EST HUMAN	EST HUMAN	N	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	EST_HUMAN	FZ	EST HUMAN	EST HUMAN	EST_HUMAN	TN.	EST_HUMAN	LN	EST HUMAN	1	EST HUMAN	EST HUMAN	NT	EST_HUMAN
Top Hit Acession No.	H75690.1	AF190338.1	U90880.1	U90880.1	W37985.1	BF510986.1		AW847284.1		N58946.1	AI016833.1	AA324242.1	AL163281.2	AA889972.1	AF128894.1	A1033980.1	AW799337.1	BF038198.1	D10548.1	A1432661.1	X68366.1	AW962164.1	11545814 NT	AI420786.1	AI420786.1		BE737895.1
Most Similar (Top) Hit BLAST E Value	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	· 6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03
Expression Signal	2.06	0.82	1.24	1.24	1.05	3.62	1.28	0.93	1.18	0.87	1.37	6.82	1.83	0.89	1.97	7.04	2.44	1.41	8.21	1.94	1.5	2.23	2.21	1.89	1.89	3.91	3.54
ORF SEQ ID NO:			23062			23311	23407	23551				24271	24758			L _	26724		27506		L	l		28356			28479
Exan SEQ ID NO:	13132	13187	13256	13256	ı	13524	13624	13757	13784	14129	14166	14485	14983	14996	16365	16468	L	16569	17297	17626	17789	18015		18102	18102		18228
Probe SEQ ID NO:	3208	3264	3336	3336	3499	3610	3711	3846	3873	4231	4267	4597	5115	6129	9059	9899	9920	6689	7509	7776	7849	8127	8186	8220	8220	8350	8351

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Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
9185	18922		1.69	6.0E-03 A	F010498.1	TN	Rhodobacter capsulatus strain SB1003, partial genome
							Methanobacterium thermoautotrophicum from bases 429192 to 450296 (section 39 of 148) of the complete
8310	19519		3.6	6.0E-03		NT	genome
9372	19038		1.34	6.0E-03	6.0E-03 AB026356.1	NT	Anguilla japonica mRNA for activin B, complete cds
8392	19583		2.21	6.0E-03	6.0E-03 U30780.1	F	Pneumocystis carinil f. sp. ratti guanine nucleotide binding protein alpha subunit (pog1) gene, complete cds
8722	19262		1.57	6.0E-03	6.0E-03 BE788019.1	EST_HUMAN	601482621F1 NIH_MGC_68 Homo sapiens cDNA clone IMAGE:3885388 5'
9741	19274		1.34	6.0E-03	6.0E-03 AJ245480.1	LN	Brassica napus sig gene for S-locus glycoprotein, cultivar T2
653	10589	20408	1.9	5.0E-03	L25105.1	TN	Chlamydia trachomatis partial ORFB; aminoacy/-tRNA synthase, complete cds; complete ORFA, and grpE- like protein, complete cds
653	10589	20407	6,1	5.0E-03	1,25105.1	TN	Ohlamydia trachomatis partial ORFB; aminoacy/-tRNA synthase, complete cds; complete ORFA, and grpE- like protein, complete cds
28	10589		2.92	5.0E-03 L	25105.1	۲N	Chlamydia trachomatis partial ORFB; aminoacyi-tRNA synthase, complete cds; complete ORFA, and grpE- like protein, complete cds
459	10589	20407	2.92	5.0E-03	L25105.1	LΝ	Chiamydia trachomatis partial ORFB; aminoacy/-tRNA synthase, complete cds; complete ORFA, and grpE- like protein, complete cds
1096	11012	ļ	1.38	5.0E-03	AJ010457.1	TN	Arabidopsis thaliana mRNA for DEAD box RNA helicase,RH3
1552	11457		0.91	5.0E-03	1138977.1	EST_HUMAN	qd79d05.x1 Scares_testis_NHT Homo sapiens cDNA clone IMAGE:1735689 3'
2648	12515		3.44		B033006.1	NT	Homo sapiens mRNA for KIAA1180 protein, partial cds
2905	12832	22629	0.95		BE266057.1	EST_HUMAN	601194796F1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:3538799 5'
3097		22820	4.08	5.0E-03		EST_HUMAN	yc81f09.s1 Soares infant brain 1NiB Homo capiens cDNA clone IMAGE:22395 3'
3118			1.75	5.0E-03	.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 3
3129		22853	1.04	5.0E-03		EST_HUMAN	y/86g02.s1 Soares breast 2NbHBst Homo sepiens cDNA clone IMAGE:155666 3'
3240	13163		1.08	6.0E-03	6.0E-03 AJ297357.1	NT	Homo sapiens partial LIMD1 gene for LIM domains containing protein 1 and KIAA0851 gene
3641	13555	23341	4.16		1.2	NT	Pseudomonas aeruginosa strain PAO1 penicillin¦binding protein 1B (ponB) gene, complete cds
3698		23396	0.83			NT	Citrus sinensis seed storage protein citrin mRNA, complete cds
3890	13801		1.38		5.0E-03 AA299675.1	EST_HUMAN	EST12218 Uterus tumar I Homo sapiens cDNA 5' end
4207	14106	23886	0.83	5.0E-03		EST_HUMAN	yu79g10.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:240066 5'
4209	13612	23386	16.0	1 EOE-03	138914.1	IN	Oltrus sinansis seed storage protein citrin mRNA, complete cds
4503	14396	24181	0.94	5.0E-03 A	J131016.1	NT	Homo saplens SCL gene locus
4619	14507	24286	2.49	5.0E-03	5.0E-03 A1752367.1	EST_HUMAN	cn15c02.x1 Normal Human Trabecular Bone Cells Homo sapiens cDNA clone NHTBC_cn15c02 random
5541			5.34	5.0E-03 P35500	P35500	SWISSPROT	SODIUM CHANNEL PROTEIN PARA (PARALYTIC PROTEIN)

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L						,	
Probe SEQ ID NO:	Exen SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
5685	15594	25895	2.43	д С	000504		PROBABLE UBIQUITIN CARBOXYL-TERMINAL HYDROLASE FAF-Y (UBIQUITIN THIOLESTERASE FAF-Y) (UBIQUITIN-SPECIFIC PROCESSING PROTEASE FAF-Y) (DEUBIQUITINATING ENZYME FAF-Y) (FAT FACETS PROTEIN RELATED, Y-LINKED) (UBIQUITIN-SPECIFIC PROTEASE 9, Y
5977	L				BE30004 4	SWISSPROI	CHROMOSOME)
609 P	15101	24877		5.0F-03	AR025024 1	ES - HOWAN	Wuset435411 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:2980871 3
6566	16424	28605		_	AB016816.1	I-N	Home centare MACI 4 TDNA
6794		26865			P48982	SWISSPROT	PETA-GAI ACTOSIDASE BEECIDES AS
7006	16883		6.92	-	M81132.1	LN	Maries complement seconds (CRS)
8103	17993		7.44		T19586.1	EST HUMAN	1994F Heart Home septems ONA clare 604
8310	18187	28435	3.17	5.0E-03	AW170334.1	EST HUMAN	xn59g05.x1 Soares_NHCeC_centreal_tumor Homo sapiens cDNA clone IMAGE:2698040 3' similar to contains L1.2. L repelitive element
8310	18187	28436	3.17	5.0E-03	4W17D334.1	EST HIMAN	xn69g06.x1 Soares_NHGeC_cervical_tumor Home saplens cDNA clone IMAGE:2698040 3' similar to
8399		28527	1.95	<u>-</u>	149153.1	EST HIMAN	Wholenger of Stretament along to Avanage 1
8433	18307	28563	1.77		10946753 NT	L	Mis misculus hasheling hasheling (#837.220) Horno sapiens cDNA clone IMAGE:70686 5'
8644	18508		3.73		3E048055 1	EST HIMAN	1990 invacional righter interest in No. 1990 (LOC58212), mRNA 1746 of A. NOI. CAMB. Brief U.
9327			5.04		Γ	LN	Gallis cellis character Lord 2 - Lord Sapiens CUNA clone IMAGE:2291622 6
9460	18091		8.65		Γ	Į.	Canas Vallus gryca arenyde-5-prosphate denydrogenase mRNA, complete cds Brinia malaxi V chromosoms
9555	19153		1.55			Z	Human pro-global type collegen (COI 204) wins award #24
9592	19174	:	1.28	5.05-03	AA456587 1	H HIMAN	ZX75a03.s1 Scares overy tumor NHOT Home explens CDNA clame IMAGE:809548 3' similar to
9816	19529		4.18		3F572332.1	1	602077774E1 NIH MGC 42 Home control OXIDASE A2 COMPONENT;
9791		25197	1.94	6.0E-03 A	_	1	UI-H-BI3-akf-608-0-11 st NCI CGAP Subs Demonstrate Control invage 4252002 6
8086			1.4	5.0E-03	Γ	1	COLLAGEN ALPHA 1/VII) CHAIN PRECLIBSOR JONG CHAIN COLLAGEN ALPHA 1/VII) CHAIN PRECLIBSOR JONG CHAIN COLLAGEN
8358	. I		1.41	5.0E-03	1668709.1	Г	20/4005.x5 Spares fetal ling NhH 10W Home colors CON 11 11:05 2000
232	10201	20015	2.97	4.0E-03 A	4.0E-03 AW 500198.1	Т	U-HF-BNO-akc-h-04-0.11 r1 NIH MGC 50 Home capitals contact that of contact the contact
318	10280	20087	2.18	4.0E-03 R46482.1		EST HUMAN	V451e04.s1 Soares infant brain 1NIB Homo sentions ONIA also 1110 Contracts
436	10380	20203	0.84	4.0E-03 P54675		Т	PHOSPHATIDYLINOSITOL 3-KINASE 3 / DIA KINASE / DITONIS 2 KINASE / CALL
288	10526	20333	3	4.0E-03 A	19.1	Τ	07/3012.81 Scares NFI T GBC St Homoscanions along March 17/2012.81
88	10785	20637	1.81	4.0E-03 R46482.1		Т	VEST604.51 Soares infant brain 1NIB Homo septem CONIA close 1NA CE conconi
88	10819		3.43	4.0E-03 A	4.0E-03 AW749101.1	Т	RC3-BT0333-110100-012-701 BT0333 Home Saviess CONA Gigle INMAGE 339888 3
1133	11047	20889	32.55	4.0E-03 A		Ī	z81a08.r1 Stratagene colon (#03770/4) Home saries a DNA
1153	11088	20910	1.68	4.0E-03 A	W794740.1	Т	RC8-UM0014-170400-023-G01 UM0014 Home satisface CDNA
					1	7	Which stands of the transfer o

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Į		1		1		7	1	1	<u> </u>]		Τ	Τ	Τ	Т	Τ	Τ	T	Τ	Τ	-		Γ.	Г	Τ	Т	Τ	Τ	Т	Т	Т	Т
	Top Hit Descriptor	zs59a01,r1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:701736 5'	AV708305 ADC Homo sapiens cDNA clone ADCAKB06 5'	Rattus norvegicus type 1 astrocyte and olfactory-limbic associated protein AT1-46 mRNA commissed constructions	281a08.r1 Stratagene colon (#937204) Homo sariens cDNA clone IMAGE:510908.5	601304161F1 NIH MGC_21 Homo sepiens cDNA clone IMAGE:3638510 5	RC6-UM0014-170400-023-G01 UM0014 Homo saplens cDNA	Homo sapiens X28 region near ALD locus containing dual specificity phosphatase 9 (DUSP9), ribosomal protein L188 (RPL188). Ca2+(Calmodulin-dependent protein kinase (CAMKI) greating transporter (CRTB)	CDM protein (CDM), adrenoleukodystrophy protein >	Homo saplens X28 region near ALD locus confaining dual specificity phosphatase 9 (DUSP9), ribosomal protein L188 (RPL18a). Ca2+(Calmodulin-dependent motein kinase (CAMKI) creating transporter (CRTB)	CDM protein (CDM), adrenoleukodystrophy protein >	Homo sapiens polyglutamine-containing C14ORF4 gene	Homo sapiens polyglutamine-containing C14ORF4 gene	Homo sapiens chromosame 21 segment HS21C084	x99604.x1 NCI_CGAP_Co18 Homo saplens cDNA clone IMAGE:2665279 3'	x98f04.x1 NCI_CGAP_Co18 Homo septens cDNA clone IMAGE:2665279 3'	OLFACTORY RECEPTOR 611 (OLFACTORY RECEPTOR-LIKE PROTEIN OLF1)	OLFACTORY RECEPTOR 511 (OLFACTORY RECEPTOR-LIKE PROTEIN OLF1)	Mus musculus tumor susceptibility protein 101 (tsg101) gene, complete cds	Homo sapiens TNNT1 gene, exons 1-11 (and joined CDS)	ab18a08.x5 Stratagene lung (#937210) Homo saplens cDNA clone IMAGE:8411423' similar to contains Alu	repetitive element;	Foot and mouth disease virus serotype A-12 119ab capsid protein VP3	Drosophila melanogaster anon2D7 (anon2D7) mRNA, complete cds	Rattus norvegicus beta-catenin binding protein niRNA, complete cds	(HPRG)	MAJOR SURFACE-LABELED TROPHOZOITE ANTIGEN PRECURSOR	Rattus norvegicus opsin gene, complete cds	601076015F1 NIH_MGC_12 Homo sapiens cDNA clone IMAGE:3461954 5'	Lycopersicon esculentum knotted 3 protein (TKn3) mRNA, complete cds	MUCIN 2 PRECURSOR (INTESTINAL MUCIN 2)	Dictyostelium discoideum AX4 development protein DG1122 (DG1122) gene, partial cds
	Top Hit Database Source	EST_HUMAN	EST_HUMAN	Ę	EST HUMAN	EST HUMAN	EST_HUMAN		NT		뒫	NT	TN	N	EST HUMAN	EST_HUMAN	SWISSPROT	SWISSPROT	TN	LN		EST_HUMAN	NT	NT	IN	SWISSPROT	SWISSPROT	NT	EST_HUMAN	Г	ISSPROT	NT
	Top Hit Acesslan No.	AA284374.1	AV708305.1	U33472.1	AA099777.1	BE410556.1	AW794740.1		U52111.2		U52111.2	AJ277365.1	AJ277365.1	AL163284.2	AW188426.1	AW188426.1	Q13606	Q13606	AF060868.1	AJ011712.1		AI732754.1	J02187.1	AF005859.1	AF169825.1	P04196	P21849	U22180.1	BE548453.1	1	Q02817	AF111944.1
	Most Similar (Top) Hit BLAST E Value	4.0E-03	4.0E-03	4.0E-03	4.0E-03	4.0E-03	4.0E-03		4.0E-03			4.0E-03	4.0E-03	4.0E-03	4.0E-03	4.0E-03	4.0E-03	4.0E-03	4.0E-03	4.0E-03				4.0E-03/	4.0E-03	4.0E-03	4.0E-03	4.0E-03				4.0E-03
	Expression Signal	1.33	1.31	22	20.87	1.66	1.46		1.89		1.89	2.86	2.86	1.02	0.94	0.94	0.8	0.83	0.85	1.79		1.13	13.17	1.68	19.53	4.23	1.5	3.23	1.76	1.52	4.14	3.61
	ORF SEQ ID NO:	21040		21480	21757		22019		22290		22291	22414	22415	22418	23207	23208	23278	23278	23556			24190	24796	24916	25039	25527	25529	<u> </u>	25827	26190	26291	26762
	Exon SEQ ID NO:	11189	11472	11610	11864	12087	12117		12400		12400	12526	12528	12530	13403	13403	'		13763	13823	,	14402	15030	15149	15236	15457	15459	15612	15714	16045	16136	16570
	Probe SEQ ID NO:	1281	1568	1709	1971	2200	2232		2526		2526	2659	2659	2864	3487	3487	3572	3836	3852	3914	į	200	9184 84	5225	5315	5540	5542	6704	6809	6062	6271	0890

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		2090013 3' similar to contains Alu					to.	ar to contains Alu repetitive	32 3' similar to contains element	ı					similar to contains Aiu repetitive							5.						,689 5'				KE7~784B0712 5'
Top Hit Descriptor	Homo sapiens KIAA0345 gene product (KIAA0345), mRNA	te49b11.x1 Soares_NFL_T_GBC_S1 Homo sablens cDNA clone IMAGE.2090013 3' similar to contains Alu repetitive element:	Homo saplens chromosome 21 segment HS21C009	Homo sapiens chromosome 21 segment HS21C078	Homo sapiens chromosome 21 segment HS21C006	PM4-BN0138-180600-002-b08 BN0138 Homo sapiens cDNA	601118164F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:3028095 5'	7q74c09.x1 NCJ_CGAP_Lu24 Homo sepiens cDNA clone IMAGE: 3' similar to contains Alu repetitive element:contains element MER31 repetitive element:	nh02c07.x1 NCI_CGAP_Kid11 Hamo saplens cDNA clone IMAGE.2953932 3' similar to contains element	LTR5 repetitive element;	RC3-ST0281-240400-015-f03 ST0281 Homo sapiens cDNA	Homo sapiens Grb2-associated binder 2 (KIAA0571), mRNA	Homo sapiens protein kinase CK2 catalytic subunit alpha gene, exon 1	Homo sapiens protein kinase CK2 catalytic subtinit alpha gene, exon 1	no73c05.s1 NOL_CGAP_Pr2 Homo sapiens cDNA clone IMAGE:782984 similar to contains Alu repetitive	element;	Homo saplens MHC class 1 region	S.cereale (cv. Halo) mRNA for triosephosphate isomerase	Mus musculus intestinal trefoil factor gene, partial cds	Mus musculus intestinal trefoli factor gene, partial cds	Arabidopsis thaliana rpoMt gene	601237982F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3609933 5'	IL2-UM0076-240300-056-D03 UM0076 Hamo sapiens cDNA	Mus musculus alpha-1(XVIII) collagen (COL18A1) gene, exon 1 and 2	C.elegans samdc gene	AV762392 MDS Homo saplens cDNA clone MDSBSG01 5'	AV762392 MDS Homo sapiens cDNA clone MDSBSG01 5'	ah04f09.y5 Gessler Wilms tumor Homo sapiens cDNA clone IMAGE:1155689 5	S.cereale (cv. Halo) mRNA for triosephosphate isomerase	Rattus norvegicus gdnf gene	xu8.P10.H3 conorm Homo saplens cDNA 3'	DKFZp781B0712 r1 761 (synonym: hamy2) Homo saplens cDNA clone DKFZp781B0712 5
Top Hit Database Source		1948			NT	EST_HUMAN PM	EST_HUMAN 601	FST HUMAN efer	✝	EST_HUMAN LTF	EST_HUMAN RC			NT		EST_HUMAN eler			NT		NT	EST_HUMAN 601	EST_HUMAN IL2-			EST_HUMAN AV		EST_HUMAN aho	NT S.c.		EST_HUMAN xu8	FST HIMAN DK
Top Hit Acession No.	7682067 NT	A1553983 1 E			AL163206.2 N	BE815173.1 E	BE298290.1	BF224125.1 E		AW614596.1 E	AW819141.1 E	11436955 NT	AF011920.1 NT	AF011920.1		AA468110.1 E	AF055066.1 NT		U46858.1	U46858.1 NT	Y09006.1	BE379296.1 E	AW802687.1 E	U34606.1 NT	Y12500.1 NT	AV762392.1 E		AI792278.1 E	Z32521.1 N		AI536141.1 E	A1 4400A7 4
Most Similar (Top) Hit BLAST E Value	4.0E-03	4 0F-03 /		4.0E-03/	4.0E-03	4.0E-03	4.0E-03	4.0E-03		4.0E-03/	4.0E-03/	4.0E-03	3.0E-03	3.0E-03		3.0E-03	3.0E-03/	3.0E-03	3.0E-03	3.0E-03	3.0E-03	3.0E-03	3.0E-03	3.0E-03	3.0E-03	3.0E-03/	3.0E-03/	3.0E-03	3.0E-03	3.0E-03/	3.0E-03	4 OF A2
Expression Signal	2.02	7.4	4.41	3.38	5.64	3.85	2.25			1.57	1.57	1.52	1.62	7.66		1.84	0.87	3.9	1.12	1.12	0.84	4.53	2.62	1.95	6.75	7.43	7.43	1.58	1.34	11.91	4.37	88.0
ORF SEQ ID NO:	26841	27123		27229	28609							25168	20148	20638		21401			22029	22030		22765	22833	23089		23591	23592	23632		23978	24096	24300
Exen SEQ ID NO:	16653	16933	L	1_	18345	19725	19009	19202	L	19840	19236	19416	10323	10787		11542	12097	12132	12133	12133	12890	12971	13037	13290	13297	13805	13805	13858	13980	14194	14311	14518
Probe SEQ ID NO:	8774	7058	7151	7169	8472	9282	3317	9628		8998	8679	8820	387	861		1638	2211	2248	2249	2249	2963	3044	3112	3371	3379	3895	3895	3950	4058	4298	4417	4630

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Single Exon Probes Expressed in Heart

				_			_	_	•							_		_		_						_	_		_	
Top Hit Descriptor	ab18a08.x5 Stratagene lung (#937210) Homo sapiens cDNA clone IMAGE:841142.3' similar to contains Alu repetitive element;	601482715F1 NIH_MGC_68 Homo sapiens cDNA clone IMAGE:3885483 5'	Homo sapiens RAP1, GTPase activating protein 1 (RAP1GA1) mRNA	Homo saplens RAP1, GTPase activating protein 1 (RAP1GA1) mRNA	Homo saplens hypothetical protein FLJ10539 (FLJ10539), mRNA	Mus musculus mRNA for hypothetical protein (ORF2 ortholog)	ea13f10.rl Soares_NhHMPu_S1 Homo saplens cDNA clone IMAGE:813163 5'	Kluyveromyces mardanus pcpl3 gene for purine-cytosine permease	Oryza sativa gene for bZIP protein, complete cds	2527b04.s1 Scares_parathyroid_tumor_NbHPA Homo sapiens cDNA clone IMAGE:304783 3'	HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN A2 HOMOLOG 1 (HNRNP A2/A))	Homo sapiens chromosome 21 segment HS21(X)68	NONSTRUCTURAL PROTEIN V	hh80f10.x1 NCI_CGAP_GU1 Homo sapiens cDNA clone IMAGE:2969131 3' similar to contains L1.11 L1	repetitive element;	Arabidopsis thallana DNA chromosome 4, contig fragment No. 85	ov03d12.x1 NCI_CGAP_Kid3 Homo sapiens cDNA clone IMAGE:1636247 3' similar to gb:X57138_rna1 HISTONE H2B.2 (HUMAN);	CIRCUMSPOROZOITE PROTEIN PRECURSOR (CS)	RETROVIRUS-RELATED POL POLYPROTEIN CONTAINS: REVERSE TRANSCRIPTASE;	ENDONUCLEASE	HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN AZ HOMOLOG 1 (HNRNP AZA))	Homo sapiens chromosome 21 segment HS21C103	Homo saplens ATP/GTP-binding protein (HEAB), mRNA	Pneumocystis carinii kexin-like serine endoprotease mRNA, partial cds	Homo saplens golgin-like protein (GLP) gene, complete cds	Homo sepiens trinucleotide repeat DNA binding protein p20-CGGBP (CGGBP) gene, complete cds	Homo sapiens trinucleotide repeat DNA binding protein p20-CGGBP (CGGBP) gene, complete cds	RETROVIRUS-RELATED POL POLYPROTEIN (CONTAINS: REVERSE TRANSCRIPTASE; ENDONUCLEASE)	promma-5.E07.r bytumor Homo saplens cDNA 8'	ot77b10.s1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA clons IMAGE:1622779 3' similar to contains L1.t3 MER26 repetitive element ;
Top Hit Database Source	EST_HUMAN	EST_HUMAN	INT	LN.	NT	NT	EST_HUMAN	۲	IN	EST_HUMAN	SWISSPROT	F.	SWISSPROT		EST_HUMAN	TN	EST HUMAN	SWISSPROT		SWISSPROT	SWISSPROT	Į.	L	NT	NT	LN	TN.	SWISSPROT	EST_HUMAN	EST HUMAN
Top Hit Acession No.	AI732754.1	BE787945.1	4506414 NT	4506414 NT	8922499 NT	AJ249981.1	AA456701.1	AJ011419.1	AB021736.1	N92580.1	P51989	AL163268.2	Q9QM81		AW613774.1	AL161589.2	A1016731.1	P08672		P11369	P51989	AL163303.2	5803028 NT	AF009222.1	AF266285.1	AF094481.1	AF094481.1	P11369	AI525056.1	AA993154.1
Most Similar (Top) Hit BLAST E Value	3.0E-03	3.0E-03	3.0E-03	3.0E-03	3.05-03	3.0E-03	3.0E-03	3.0E-03		3.05-03	3.0E-03	3.0E-03			3.0E-03	3.0E-03	3.0E-03							3.0E-03		3.0E-03	3.0E-03	3.0E-03	3.05-03	3.0E-03
Expression Signal	1.54	60,7	1.11	1.11	3.17	1.41	9.97	1.54	3.71	2.01	1.29	1.57	1.38		8.52	4.1	8.8	3.48		1.29	1.46	4.05	2.69	2.11	2.01	3.06	3.06	1.93	2.02	1.54
ORF SEQ ID NO:	24399	24416	24733	24734	24837	25380	26974	26297	26468	26809		27011			_	27297	27317	L				27831					28878	28942		28780
Exan SEQ ID NO:	14613	14630	14959	14959		15330	15851	18141	16303	16620	16806	16818	16868		17076	17107	17121	I_{-}	1	╛	17592	17686	1818 1818			18590	18590	18654	19542	1 1
Probe SEQ ID NO:	4727	4745	5089	6809	5220	5410	6946	429	6442	6741	6928	6940	6991		7199	7230	7244	7598		8	7742	7836	8222	8685	8744	8773	8773	8841	8906	9103

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					5	יסיי רוויסאם פופי	Cingle Lyon Flores Capiessed in Hear
Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
9159			1.35	3.0E-03	3.0E-03 AB009668.1	IN	Homo sapiens gane for CMP-N acetylaenramining acid hudrondoor
88		25296	2.33	3.0E-03	3.0E-03 AJ296282.1	Ł	Reffus norveoleus mRNA for connoving (2008)
504	_ [0.89	2.0E-03 Q04652	Q04652	SWISSPROT	RING CANAL PROTEIN (MELCIL DEDICE)
504	- 1	20259	0.89	2.0E-03 Q04652	Q04652	SWISSPROT	RING CANAL PROTEIN (VELCH PROTEIN)
768	12677		7.01	2.0E-03 T	T70874.1	FST HIMAN	MARKS ALVOIDEM (NELCOT PROTEIN)
1340			1.93	2.0E-03		LN TN	Mirran glybe 2 placen intel spieen 1NPLS Home sapiens cDNA clone IMAGE:108341 5'
1343	11249	21106	1.59	2.0E-03	AA661605.1	EST HUMAN	PURBETO 15 NOT COAD AND Home seemed and 1
1353	11259	21115	12.6	2.0E-03	2.0E-03 AF284446.1	N.	Homo seplens trimor-related protein DDC2 (DDC2)
1474	11379	21243	1.11	2.0E-03 P.	P48509	SWISSPROT	PLATELET-ENDOTHELIAL TETRASPAN ANTIGEN 3 (PETA-3) (GP27) (MEMBRANE GLYCOPROTEIN SFA-1) (CD161 ANTIGEN)
1500	11404	21263	1.91	2.0E-03	4557836 NT	Ę	Homo septens procollagen-lysine, 2-oxoglutarate 5-dloxygenese (lysine hydroxylase, Ehlers-Danlos syndrome type VI) (PLOD) mRNA
1500	11404	21264	1.91	2.0E-03	4557836 NT		Homo sapiens procellagen-lysine, 2-oxoglutarate 5-dloxygenase (lysine hydroxylase, Ehlers-Danlos syndrome
1573	11477		4.94	2.0E-03 P29400	1	ISSPROT	THE TABLE STANDARDED TO SEE THE STANDARD STANDAR
1733	11634	21502	1.38	2.0E-03 /	VA450138.1	Т	242910 11 Sparse Intoles Annie Includes Annie Includes Annie Include Ann
1950		21731	2.17	2.0E-03 /	2.0E-03 AF302691,1	Т	Mus musculus movelin accession forter 3 III.
2203	_1	21992	1.04	2.0E-03 AL			Homo saplens chromosome 34 second Listed Auto Auto
2535			4.62	2.0E-03 A		F HIMAN	III-H-RICadio-10-01 of NCI OCAB 6:15-11
3370	- 1	23088	4.03	2.0E-03 A	Γ	Т	24/2910 r1 Soares total faths Nhoules on London Sapiens culva clone IMAGE:2717010 3
3376	13294	23093	1.02	2.0E-03 E	2.0E-03 BF568955.1	Т	602183960T1 NIH MGC 42 Home canifors all March 11 March 20 Home canifors all March 20 Home canifors al
3615	13529	23315	4.87	2.0E-03 X87344.1			H. saplens DMA, DMB, HLA-Z1, IPP2, LMP2, TAP1, LMP7, TAP2, DOB, DOB2 and RING8, 9, 13 and 14
6986 8	13780	23572	0.92	2.0E-03 A	1		Rattus noveolcus mRNA for SREB1 complete at
4024	13927	23703	2.15	2.0E-03 P03374		/ISSPROT	ENV POLYPROTEIN CONTAINS: COAT DECATE AND ASSESSMENT OF THE ASSESSMENT ASSESS
4127	14027		9.77	2.0E-03 U68491.1			Rattus novemieres Entrementaries Control Entre Graz, COA I PROTEIN GP38]
446	14340	24130	1.93	20E-03 L			Drosoohila malanoaster shortsichted class 2 (Alic) - Daile
4448	14340	24131	1.93	2.0E-03 L42512.1			Drosophila melanogaster shortsighted class 2 (shs) mRNA complete cds
4608	14498	24285	0.94	2 OF-03	2 DE-03 A F223301 4	1	Homo seplens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively
4612	14500		1.7	2.0E-03 R		HIMAN	opinical model to the second of the second o
2077	14947		0.82	2.0E-03 A.	2.0E-03 AJ245167 1		Camplis dramadating authority and 2000 from Sapiens CDNA clone IMAGE:180890 3'
5194	15057	24820	1.04	2.0E-03 A	Ī		Homo senions extracell for altrophetic location.
5194	15057	24821	1.04	2.0E-03 A		Ę	Homo saplens extracellular olycomorphin length precursor, gene, complete cds
					1		Complete Commenter Strongly National Processor, gene, complete cds

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Table 4
Single Exon Probes Expressed in Heart

Droha	Evon			Most Similar		ij	
SEQ ID	SEO ID	ORF SEQ ID NO:	Expression Signal	(Top) Hit BLAST E Value	Top Hit Acession No.	Database Source	Top Hit Descriptor
6372	15292	25128	1.37	2.0E-03	BF241410.1	EST_HUMAN	601876385F1 NIH_MGC_55 Homo sapiens cDNA clone IMAGE:4104692 5'
5457	19445	25438	1.78	2.0E-03	AB014593.1	IN	Homo sapiens mRNA for KIAA0693 protein, pertial cds
5501	15420	25482	1.87	2.0E-03	U63711.1	NT.	Xenapus Iaevis xefitim mRNA, complete cds
5718		25725	3.25	2.0E-03	P23477	SWISSPROT	ATP-DEPENDENT NUCLEASE SUBUNIT B
5718			3.25		P23477	SWISSPROT	ATP-DEPENDENT NUCLEASE SUBUNIT B
5829	15735	25846	1.82	2.0E-03	Q95203	SWISSPROT	CARBONIC ANHYDRASE-RELATED PROTEIN 2 PRECURSOR (CARP 2) (CA-RP II) (CA-XI)
5829	15735	25847	1.82	2.0E-03	Q95203	SWISSPROT	CARBONIC ANHYDRASE-RELATED PROTEIN 2 PRECURSOR (CARP 2) (CA-RP II) (CA-XI)
5831	15737	25849	7.14	2.0E-03	BF308187.1	EST_HUMAN	6018B7434F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:4121408 5
5850	15758	25874	2.2	2.0E-03	Q9UKP4	SWISSPROT	ADAM-TS 7 PRECURSOR (A DISINTEGRIN AND METALLOPROTEINASE WITH THROMBOSPONDIN MOTIFS 7) (ADAMTS-7) (ADAM-TS7)
5861	'		1.95	2.0E-03	X94451.1	N _T	L.esculentum mRNA for lysyl-IRNA synthetase (LysRS)
5978	15884		1.36	2.0E-03	Al991089.1	EST HUMAN	wu36h09.x1 Soares_Dieckgraefe_colon_NHCD Homo sapiens cDNA clone IMAGE:2522177 3' similar to SW:RL29_HUMAN P47814 60S RIBOSOMAL PROTEIN L29 ;contains element MSR1 repetitive element ;
6212	15994	26129	2.88	2.0E-03	BE067986.1	EST_HUMAN	CM4-BT0366-061299-054-d01 BT0366 Homo sapiens cDNA
6691	16571	26763	2.11	2.0E-03	AW 692004.1	EST_HUMAN	hf37b08.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA done IMAGE:2834035 3' similar to TR:Q60976 Q60976 JERKY.;
6784	16663	26853	5.19	2.0E-03	N20287.1	EST_HUMAN	yx42g06.s1 Soares melanocyte 2NbHM Homo sapiens cDNA clone IMAGE:264442.3' similar to contains L1.b2 L1 repetitive element;
6784	16663	26854	5.19	2.0E-03	N20287.1	EST_HUMAN	yx42g08.s1 Soares melanocyte 2NbHM Homo sapiens cDNA clone IMAGE:264442.3' similar to contains L1.b2 L1 repetitive element;
7513	17301	27508	3.12	2.0E-03	P24821	SWISSPROT	TENASCIN PRECURSOR (TN) (HEXABRACHION) (CYTOTACTIN) (NEURONECTIN) (GMEM) (JI) (MIOTENDINOUS ANTIGEN) (GLIOMA-ASSOCIATED-EXTRACELLULAR MATRIX ANTIGEN) (GP 150- 226) (TENASCIN-C) (TN-C)
7775	11		5.41	•		EST_HUMAN	2s10a08.s1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:084754 3'
8379	18256		2.78	2.0E-03		NT	Human dystrophin gene
8779	18350	26520	1.78	2.0E-03	P07354	SWISSPROT	PROTEOGLYCAN LINK PROTEIN PRECURSOR (CARTILAGE LINK PROTEIN) (LP)
8829	18642		1.77	2.0E-03	BF330909.1	EST_HUMAN	RC3-BT0333-310800-115-g04 BT0333 Homo sapiens cDNA
8836	18649	98638	12.17	2.0E-03	211740.1	NT	H.sapiens variable number tandem repeat (VNTR) locus DNA
8052	18835		2.07	2.0E-03	AJ625745.1	EST_HUMAN	ty65h03.x1 NCI_CGAP_Kid11 Homo sepiens cDNA clone IMAGE:2283989 3' similer to SW:VATG_MANSE Q25532 VACUOLAR ATP SYNTHASE SUBUNIT G;
9086		29115		2.0E-03		NT.	Homo sapiens SEL1L (SEL1L) gene, partial cds
9113			4.48	2:0E-03		NT	Camelus dromedarius cyhp19 gene for immunoglobulin heavy chain variable region
9322	19703		2.78	2.0E-03	AV697866.1	EST_HUMAN	AV697968 GKC Hamo sapiens cDNA clone GKCGXD05 5'

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Page Expression Page P
SEQ ID DNO: Signal Most Similar Top Hit Acession Top Hit Top Hit Acession No: Signal No: Signal No: Signal No: Signal No: Signal No: Signal No: Source No: Source 15086 2.0E-03 Y00508.1 NT NT NT NT NT NT NT N
Exon NO: CRF SEQ Signal Signal 19066 Expression Signal Velue 19066 Most Similar Signal Velue 19060 Top Hit Acession Velue 2.0E-03 Top Hit Acession No. Top Hit Acession Source Source 1.59 Top Hit Acession Velue 2.0E-03 Top Hit Acession Velue 2.0E-03 Top Hit Acession No. Top Hit Acession Source 2.0E-03 Top Hit Acession No. Top Hit Acession Source 1.59 Top Hit Acession 2.0E-03 Top Hit Acession No. Top Hit Acession Source 2.0E-03 No. Source Source 3.0E-03 Alt Rid Source Alt Rid Source 3.0E-03 Alt Rid Source Alt Rid Source 3.0E-03 No. Source 3.0E-03 No. Source 3.0E-03 Alt Rid Source Alt Rid Source 3.0E-03 No. Alt Rid Source 3.0E-03 Alt Rid S
Exon NO: ORF SEQ Signal Signal 19085 Expression Signal No: Most Similar Signal No: Top Hit Signal Signal No: Top Hit Signal No:
Exon NO: ORF SEQ Signal Signal 19087 Most Similar Signal Signal 19087 Top Hit Acession Signal PLAST E 159 Top Hit Acession Signal PLAST E 159 Top Hit Acession Signal PLAST E 159 Top Hit Acession Post E 2 0E-03 N0508.1 Top Hit Acession Source NT 19087 2.62 2.0E-03 Y00508.1 NT 19569 2.67 2.0E-03 Al76037.1 EST HUMAN 19571 2.016 3.62 2.0E-03 Al76037.1 EST HUMAN 10740 20587 2.38 1.0E-03 Al720263.1 EST HUMAN 10740 20588 2.38 1.0E-03 Al720263.1 EST HUMAN 10740 20586 2.38 1.0E-03 Al720263.1 EST HUMAN 11063 20806 1.44 1.0E-03 Al865788.1 EST HUMAN 11063 20806 1.44 1.0E-03 Al86578.1 EST HUMAN 11063 20806 1.44 1.0E-03 Al86578.1 EST HUMAN 11075 2186 8.02 1.0E-03 Al86578.1 EST HUMAN 11075 2186 8.02 1.0E-03 Al86578.1 EST HUMAN 13079 22876
Exon NO: 19066 CRF SEQ 25278 Expression Signal 1.59 (Top) Hit LASTE 1.59 Top Hit Acession Value 2.0E-03 Top Hit Acession No: 19087 Top Hit Acession Value 1.59 Top Hit Acession 2.0E-03 Top Hit Acession No: 1.59 Top Hit Acession Value 2.0E-03 Top Hit Acession No: 1.59 Top Hit Acession 2.0E-03 Top Hit Acession No: 1.0E-03 Top Hit A
Exon NO: ORF SEQ Signal 19068 Expression Signal 10068 (Top) Hit Signal 1.59 Top Hit LASTE Value 2.02 Top Hit Source 3.02 Hit Source 3.02 Top Hit Source 3.02 Top Hit Source 3.02 Hit Source 3.02 Hit Source 3.02 Top Hit Source 3.02 Top Hit Source 3.02 Top Hit Source 3.02 Top Hit Source 3.02 Top Hit Source 3.02 Hit Source 3.02 Hit Source 3.02 Hit Source 3.02 Hit Source 3.02 Hit Source 3.02 Hit Source 3.02 Hit Source 3.02 Hit Source 3.02 Hit Source 3.02 Hit Source 3.02 Hit Source 3.02 Hit So
Exon NO: 19068 ORF SEQ 25278 Expression Signal 1.59 (Top) Hit LASTE Value 2.02 Top Hit 2.02 Top Hit 2.047 Top Hit Pathess NT Top Hit Source Source Source NT 19088 25278 1.59 2.0E-03 M-163203.2 NT 19134 2.02 2.0E-03 M-163203.2 NT 19521 1.57 2.0E-03 M-163203.2 NT 19522 1.57 2.0E-03 M-163203.2 NT 10740 20587 2.38 1.0E-03 M-169768.1 NT 10740 20587 2.38 1.0E-03 M-169768.1 EST_HUMAN 10740 20588 2.38 1.0E-03 M-169768.1 EST_HUMAN 11015 20508 1.0E-03 M-169477.1 EST_HUMAN 11015 20508 1.44 1.0E-03 M-169477.1 EST_HUMAN 11015 20508 1.45 1.0E-03 M-1692618.1 EST_HUMAN 11015 20508 1.45 1.0E-03 M-1692618.1 EST_HUMAN 11876 21768 8.02 1.0E-03 M-1692618.1 SWISSPROT 13079 22876
Exon ORF SEQ Expression Signal (Top) Hit Acession No. Top Hit Acession Signal Top Hit Acession Signal Top Hit Acession Signal Top Hit Acession No. Top Hit Acession Signal Top Hit Acession No. Top Hit Acession Signal Top Hit Acession No.
Exon ORF SEQ Expression Signal (Top) Hit Top Hit Acession No. Top Hit Soutzee 19068 25278 1.59 2.0E-03 Y00508.1 NT 19087 9.82 2.0E-03 P4278.1 NT 19747 2.82 2.0E-03 P4278.1 NT 19569 1.57 2.0E-03 AL163203.2 NT 1977 2.0E-03 AL163203.2 NT 1977 2.0E-03 AL163203.1 EST_HUMAN 10377 20198 1.38 1.0E-03 AV697866.1 EST_HUMAN 10740 20587 2.38 1.0E-03 AI720263.1 EST_HUMAN 10740 20588 2.38 1.0E-03 AI720263.1 EST_HUMAN 11045 20806 2.38 1.0E-03 AI865788.1 EST_HUMAN 11045 20806 1.44 1.0E-03 AI865788.1 EST_HUMAN 11045 2.10E-03 AI892618.1 EST_HUMAN 11046 2.286 1.0E-03 AI892618.1 EST_HUMAN 11876 2.10E-03 AI892618.1 EST_HUMAN 11876 2.10E-03 AI892618.1
Exon ORF SEQ ID Expression ID NO: Signal Most Similar Top Hit Acession No. Signal BLAST E No. Signal 159 Most Similar Logo Top Hit Acession Signal No. Detabase Source Signal No. Signal 159 Top Hit Acession No. Signal
Exon ORF SEQ ID Expression ID NO: Signal Top Hit Acession No: Signal Most Similar Top Hit Acession No: Signal No: Signal No: Signal No: Signal No: Signal No: Signal No: Signal No: No: Signal No: No: No: No: No: No: No: No: No: No:
Exon ORF SEQ ID Expression ID NO: Signal Top Hit Acession No: Signal Most Similar Top Hit Acession No: Signal No: Signal No: Signal No: Signal No: Signal No: Signal No: Signal No: No: Signal No: No: No: No: No: No: No: No: No: No:
Exon ORF SEQ Expression No. Most Similar Top Hit Acession Buaste Signal Most Similar Top Hit Acession No. Top Hit Acession Signal No. Top Hit Acession Signal No. Top Hit Acession Signal No. Top Hit Acession Signal No. Top Hit Acession No. Top Hit Acession Signal No. Top Hit Acession No. Top Hit Acession Signal No. Top Hit Acession No.<
Exon No: SEQ ID Most Similar (Top) Hit Acession ID No. Signal No. Sign
Exon ORF SEQ ID NO: Expression Signal Most Similar PLASTE PLASTE Value Top Hit No. Top Hit Soutze Soutze Soutze Soutze Soutze 19066 25278 1.59 2.0E-03 Y00508.1 NT 19087 9.82 2.0E-03 Y00508.1 NT 19087 9.82 2.0E-03 Y00508.1 NT 19547 1.57 2.0E-03 AL163203.2 NT 19521 1.57 2.0E-03 AL163203.1 EST_HUMAN 10740 20587 2.38 1.0E-03 AI20263.1 EST_HUMAN 10740 20588 2.38 1.0E-03 AI20263.1 EST_HUMAN 10740 20588 2.38 1.0E-03 AI20263.1 EST_HUMAN 10895 20836 2.14 1.0E-03 AI866788.1 EST_HUMAN 10015 20836 2.14 1.0E-03 AI20268.1 EST_HUMAN 1015 20836 2.0E-03 AI20268.1 EST_HUMAN 10740 20836 2.14 1.0E-03
Exon ORF SEQ ID NO: Expression Signal Most Similar (Top) Hit BLASTE Top Hit No. Top Hit Soutze 19066 25278 1.59 2.0E-03 Y00508.1 NT 19087 9.62 2.0E-03 Y00508.1 NT 19186 2.92 2.0E-03 Y00508.1 NT 19547 1.57 2.0E-03 AL163203.2 NT 19521 1.57 2.0E-03 AL163203.1 EST_HUMAN 10740 20587 2.38 1.0E-03 AI720263.1 EST_HUMAN 10740 20588 2.38 1.0E-03 AI720263.1 EST_HUMAN 10740 20588 2.38 1.0E-03 AI720263.1 EST_HUMAN 10895 20836 2.14 1.0E-03 AI720263.1 EST_HUMAN
Exon ORF SEQ ID ID NO: Expression Signal (Top) Hit Top Hit Acession No. Top Hit Acession Signal No. Top Hit Acession Signal No. Top Hit Acession No. Top Hit
Exon ORF SEQ Expression NO: Most Similar Signal NO: Most Signal NO: Most Signal NO: Top Hit Acession No: Database Source No: Database Source No: Database Source No: Database Source No: Database No
Exon ORF SEQ Expression ID NO: Crop) Hit Signal Top Hit Acession No: Top Hit Acession Signal Top Hit Acession No: Top
Exon ORF SEQ Expression ID NO: Cop Hit Signal Signal Top Hit Acession No: Top Hit Acession Signal BLASTE Top Hit Acession No:
Exon ORF SEQ Expression In Inc. Top Hit Acession Signal Top Hit Acession No. Top Hit Acession No. Top Hit Acession Signal PLASTE Top Hit Acession No.
Exon ORF SEQ Expression (Top) Hit Top Hit Acession Signal Most Similar Top Hit Acession Signal Top Hit Acession Detabase No. Top Hit Acession Detabase Source Signal Top Hit Acession No. Top Hit Acession Detabase Source So
Exon ORF SEQ ID NO: Expression ID NO: Top Hit Acession Signal PLASTE Top Hit Acession No. Top Hit Acession Detablese Source 19066 25278 1.59 20E-03 Y00508.1 NT 19134 2.962 2.0E-03 Y00508.1 NT 19569 2.82 2.0E-03 A1163203.2 NT 18569 1.57 2.0E-03 A1376037.1 EST HUMAN
Exon ORF SEQ ID NO: Expression ID NO: Top Hit Acession Signal ID NO: Top Hit Acession Signal ID NO: Top Hit Acession Detablese No. Top Hit Acession Detablese No. Top Hit Acession ID NO: Detablese Source ID NO: Source ID NO:
Exon ORF SEQ Expression ID NO: Columnary Signal Top Hit Acession ID NO: Top Hit Acession ID Ac
Exon ORF SEQ Expression ID NO: Top Hit Acession Signal Top Hit Acession Database Source No. Source Source Source 19068 25278 1.59 2.0E-03 Y00508.1 NT H. sepiens M1 gene for muscarinic acetylchol
Exon ORF SEQ Expression (Top) Hit Acession ID NO: Signal BLAST E No. Soutce Soutce Set 159 2 0 E-03 Y00 FR 1 19068
Exon ORF SEQ Expression (Top) Hit Top Hit Acession Signal BLASTE No: Source Source

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					,		
Probe SEQ ID NO:	Exon SEQ ID NO:	ς S Ω	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
4717	14603		1.78	1.0E-03	AI073485.1	EST_HUMAN	ov45c04.x1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:16402623'
4717	14603	24389		1.0E-03	A1073485.1	EST_HUMAN	ov45c04.x1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:16402623'
4718	14604		6.23	1.0E-03	BE154067.1	EST_HUMAN	PM0-HT0339-200400-010-D02 HT0339 Homo sapiens cDNA
5009	14883	24849	16.98	1.0E-03	046409	SWISSPROT	APOLIPOPROTEIN A-IV PRECURSOR (APO-AIV)
5103	14971	24747	0.93	1.0E-03	AV685870.1	EST_HUMAN	AV685870 GKC Homo saplens cDNA clone GKCDME11 5'
6250	15173	24946	2.03	1.0E-03	AA280951.1	EST_HUMAN	zs44f01.r1 NCI_CGAP_GCB1 Homo saplens cDNA clone IMAGE:700345 5'
5317	16238	25042		1.0E-03	AJ006345.1	⊥N	Homo saplens KVLQT1 gene
5349	15269	25098	2.01	1.0E-03	K03332.1	NT	Epstein-Barr virus (AG876 isolate) U2-IR2 domisin encoding nuclear protein EBNA2, complete cds
5349	15269	25097	2.01	1.0E-03	K03332.1	TN	Epstein-Barr virus (AG876 isolate) U2-IR2 domain encoding nuclear protein EBNA2, complete cds
5423	15344	25397	1.69	1.0E-03	Q02388	SWISSPROT	COLLAGEN ALPHA 1(VII) CHAIN PRECURSOR (LONG-CHAIN COLLAGEN) (LC COLLAGEN)
9999			4.08	1.0E-03	X07699.1	LN	Mouse nucleofin gene
5758	15668		8.56	1.0E-03	11526176 NT	TN	Homo saplens T-cell lymphoma invasion and metastasis 1 (TIAM1), mRNA
5824	15730	25842	1.32	1.0E-03	T87761.1	EST_HUMAN	yd93a11.r1 Soares fetal liver spleen 1NFLS Homo saplens cDNA clone IMAGE:115772 5'
5857	15763		1.4	1.0E-03	AW902585.1	EST_HUMAN	QV3-NN1024-260400-171-g05 NN1024 Homo sapiens cDNA
6242	16108	26259	2.4	1.0E-03	D16826.1	TN	Human gene for fourth somatostatin receptor subtype
6419	16281		1.37	1.0E-03	AJ229042.1	TN	Homo saplens 959 kb contig between AML1 and CBR1 on chromosome 21q22, segment 2/3
							Homo sapiens X28 region near ALD locus containing dual specificity phosphatase 9 (DUSP9), ribosomal
							protein L18a (RPL18a), Ca2+/Calmodulin-dependent protein kinase I (CAMKI), creatine transporter (CRTR),
6503	16362	28537	1.63	1.0E-03	U52111.2	NT	CDM protein (CDM), adrenoleukodystrophy protein >
8238	16394	28573	3.13	1.0E-03	M63376.1	TN	Human TRPM-2 protein gene, exons 1,2 and 3
9836	16516		5:32	1.0E-03	AJ251973.1	⊥N	Homo saplens partial steerin-1 gene
6798	16677	26867	2.39	1.0E-03	AF153980.1	NT	Homo sapiens exostoses-like protein 1 (EXTL1) gene, exons 2 through 11, and complete cds
7176	L		1.4	1.0E-03	Y11204.1	NT	V.carteri gene encoding volvoxopsin
7273	17150		4.27	1.0E-03	M30471.1	LΝ	Human class III alcohol dehydrogenase (ADH6) chi subunit mRNA, complete cds
7273	17150	27346		1.0E-03	M30471.1	±N	Human class III alcohol dehydrogenase (ADH5) chi subunit mRNA, complete cds
7541			1.85	1.0E-03	AF011400.1	IN	Thermotoga neapolitana alpha-1,6-galactosidase (aglA) gene, complete cds
7541	17392	27603	1.85	1.0E-03	AF011400.1	TN	Thermotoga neapolitana alpha-1,6-galactosidase (aglA) gene, complete cds
8056	17947	28196		1.0E-03	AW362393.1	EST HUMAN	RC1-CT0279-181099-011-a09 CT0279 Homo sapiens cDNA
8056	3 17947	L	22.37	1.0E-03	AW362393.1	EST_HUMAN	RC1-CT0279-181099-011-a09 CT0279 Homo saplens cDNA
8133	18021	28268	2.44	1.0E-03	BE170859.1	EST_HUMAN	QV3-HT0543-220300-130-a03 HT0543 Homo sapiens cDNA
_	Į						tf73e12.x1 NCI_CGAP_HSC3 Homo sapiens cDNA clone IMAGE:2246446 3' similar to TR:Q26195 Q26195
8188	ļ		2.69		AI583847.1	EST_HUMAN	PVA1 GENE:
8497	- 1				AV759949.1	EST_HUMAN	AV759949 MDS Homo sapiens cDNA clone MDSDDF11 5
9046	18831	29113	2.82	1.0E-03	BE894488.1	EST_HUMAN	601433087F1 NIH_MGC_72 Homo sapiens cDNA clone IMAGE:3918524 5'

Page 137 of 413 Table 4 Single Exon Probes Expressed in

Prode Exam OPF SEQ Expression Prode Expressed in Heart			1										1				-1	7	_	_	_	_	_	_	$\overline{}$		_	_		_	_					
Exan ORF SEQ Expression (Top Hit Acesolon Signal 19 No. Signal 19 No	oes Expressed in Heart	Top Hit Descriptor	Raffus noveoleus transformation related assets: 69 (T. 00).	te05h11.x1 NCI_CGAP_Co16 Homo saplens cDNA clone IMAGE:2063013.3' similar to contains Alu	repetitive element;	00.14088/8F1 NIH_MGC_67 Homo sapiens cDNA clone IMAGE:3872035 5	Applicable Architecture of the Application of the A	Glorables of the Care Care (APC-AIV)	Visuate mPNA 62-7359	MADAD 4 Soon feet in	1 INE-1 BEVERSE TRANSPORTED AND TO SELECT TRANSPORTED TO THE PARTY OF	Homo captons, with a martin (DED)	72/4/10 of Scarce feld. For the first interest and the felds	TEXTORS SOLD THE TRANSPORT OF THE TRANSP	Home smiles CVB47 202 Fig. 1	Transcaptions of Fig. 9 and	norno sapiens priori protein (PrP) gene, complete eds	nomo sapiens chromosome 21 segment HS21C010	nomo sapiens chromosome X open reading frame 6 (CXORF6) mRNA	#93500 x1 Soares_NSF_F8_9W_OT_PA_P_\$1 Home sapiens cDNA clone IMAGE:2367209 31	L44L) and FTP3 (FTP3) genes complete of the support	SC284072 normalized Infant brain CONTAIN	CM1-RT0614.410300-142-142-142-142-142-142-142-142-142-142	of 3c08 of Soares Infant brein 1NIB Dome	Tomo sapiens Retine derived POLI domain forth 4 / PDE 22	VISSILIA NOI CGAP Kido Home control (RPF-1), MRNA	domo saniens ensilon.1 necessidades (10) 1-153	John Sabiens ension-1 pseudogene (ICHEPT) (Jahe, 6 Tienking region	forms smith the complete control of the control of	UCOSE TRANSPORTED TYPE F 2001 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	KEZNERAMONA A BB (MINISTER)	C2-RN0120, 250400, 012 141 BN0430 11	Ome sapient 050 kh confidence and a sapient co	1-H-Blo-gab-e-09-0-Lil st NCI CGAP Substitution of the Capture of	UCLEOSIDE DIPHOSPHATE KINASE B (NDK B) (NDP KINASE BY/NM23 M3/1048)	(סו בו) (אוואניסיואול) (בי בייניי בייניי בייניי בייניי בייניי בייניי בייניי בייניי בייניי בייניי בייניי בייניי
Exam ORF SEQ Expression Signal (Top) Hit Top Hit Acess No. Top Hit Acess No. 19108 25287 1.95 1.0E-03 9507 19707 24904 4.2 1.0E-03 BET80572.1 14998 24769 0.81 9.0E-04 L11610.1 14504 2450 4.2 1.0E-03 BET80572.1 14598 24769 0.81 9.0E-04 L11610.1 14598 24769 0.81 9.0E-04 L1610.0.1 14598 24769 0.81 9.0E-04 L1610.0.1 14542 2456 1.27 8.0E-04 L1610.0.1 13989 4.2 1.0E-03 BE4037203.1 14542 24332 2.16 8.0E-04 L1610.0.1 14542 24332 2.39 8.0E-04 L16320.0 14542 24332 2.39 8.0E-04 L4855.1 15848 22437 3.26 7.0E-04 L41825.1 15876 22437 3.26 7.	gle Exon Pro		L	i i	EST HUMAN	ES L'AUMAIN	TOGGSSIMS	TA LE	Į.	EST HIMAN	SWISSPROT	5	Т	Т	NO.							HUMAN	HUMAN	HUMAN		T HUMAN				SSPROT	Т	Т		Г		
Exam ORF SEQ Expression (Trigonal Incident) Mossignal Expression (Trigonal Incident) Mossignal Expression (Trigonal Incident) Mossignal Expression (Trigonal Incident) Mossignal Expression (Trigonal Incident) Mossignal Expression (Trigonal Incident) Mossignal Expression (Trigonal Incident) Mossignal Expression (Trigonal Incident) Mossignal Incident Mossignal		Top Hit Acession No.		A 13 4 7 2 E C	RF780572 4	11910 1	P08727	AB037203 1	X96469.1	307008.1	208547			Ţ				25470	2						6005855						T			Γ		
Exan NO: ORF SEQ ID NO: Express Signe Signe 19707 Express 24287 19687 25287 Signe Signe 14998 24769 14998 24769 24769 15404 24769 24769 15764 23548 2727 13764 23548 22127 14542 24332 22127 14542 22437 3 18683 22127 3 12230 22437 3 12547 22437 3 12548 22856 1 15828 28873 2 16863 23669 1 13805 23680 0 13905 23681 0 13905 23681 0 14569 28861 2 14669 28950 6 14669 28950 6		Most Similar (Top) Hit BLAST E Value	1.0E-03	1 05 03	1 OF-03	9.0E-04	9.0E-04	9.0E-04	8.0E-04	8.0E-04	8.0E-04	8.0E-04	8.0E-04/	8.0E-04/	7.0E-04										7.0E-04		6.0E-04 K	8.0E-04 K	6.0E-04 U	8.0E-04 P	6.0E-04 A					
Exan ORI NO: 19108 19687 19108 19687 19707 1989 1989 1989 1989 1989 1989 1989 198		Expression Signal	1.95	2.15	4.2	0.81	1.58	1.42	1.39	1.27	42	2.39	2.08	2.16	1,61	1.13	3.28	1.03	2.24		3.2	2.61	3.29	2.72	3.28	1.34	0.85	0.85	3.28	4.35	3.13	2.24	2.07	6.32	2.45	
0					L					23548		24332			21562	22127	22437	22965				28973				23576	23680	23681	23769			27887	28881	28950		
Probe SEQ ID NO: NO: NO: NO: NO: NO: NO: NO: NO: NO:			Ц			L						_1	18358	18483	11684	12230	12547	13166	15828		18683	18683	19165	19300	18821	13788	13905	13905	13992	16470	17583	17650	18593	18664	18714	
		Probe SEQ ID NO:	9497	9517	9626	5131	5485	7561	1472	3843	4089	4656	8485	8816	1786	2350	2682	3243	6923	3	3	8874	9558	9780	9813	3877	3330	3889	4092	9630	7733	7800	8776	8852	8908	

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Single Exon Probes Expressed in Heart	Ton Hit Descriptor	lordings)	Т	T	7	٦	7		Т	Bos taurus micromolar calcium activated neutral professe 1 (CADMA)	zo33b08.r1 Stratagene colon (#937204) Homb sapiens cDNA close IMAGE space ri	Gonilla garilla Involucrin gene medium allele, complete cds	do 13/16/X1 Soares placenta_8to9weeks_2NbHP8to9W Homo sepiens cDNA clone IMAGE:17236193' similar to gb:X51602_cds1 VASCULAR ENDOTHE IAI GROWTHE SACTOR PROPERTY	(HUMAN);contains Alu repetitive element:	gj56h03.s1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE-1304357.9			Bos taurus micromolar calcium pertinated months in the calcium pertinated months in the calcium pertinated months in the calculation and calcu	Inf15h02.s1 NCI CGAP Pr1 Home series and Alfa	xm77h09.x1 NCI_CGAP_Kid11 Homo saniens cDNA clone IMAGE:913875	601876534F1 NIH_MGC_55 Homo sapiens cDNA clone IMAGE: 4104807 g:	Haemophilus influenzae Rd section 63 of 163 of the complete genome	O13825 AU-BINDING PROTEINENOYI-COA HYDRATASE	as70b08.x1 Barstead colon HPLRB7 Homo sapiens cDNA clone IMA CE: 222 Appa 21	O13825 AU-BINDING PROTEIN/ENOYL-COA HYDRATASE	RC3-CT0254-130100-023-f01 CT0254 Homo sapiens cDNA	Homo sapiens chromosome 21 segment HS21C078	DKFZp434D059_r1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZzz434D0Eo E	SEMICIN-2 (SILK GUM PROTEIN 2)	Homo sapiens neuropilin 2 (NRP2) gene, complete cds, alternatively spliced	nh10a10.s1 NCI_CGAP_Co1 Homo saplens cDNA clone IMAGE:951930 3' similar to gb:M21121 T-CELL SPECIFIC RANTES PROTEIN PRECIDENCE ALIMANN	(JORGAN)
gle Exon Pr	Top Hit Database	Source	100	ES! HUMAN	IN COUNTY	OWISSPRO	EST HUMAN	EST HUMAN		Ę	EST_HUMAN	Z		ESI HOMAN	EST_HOMAN	ES HUMAN	EST_HUMAN	L L	ST_HUMAN	EST_HUMAN	EST_HUMAN	Z	EST HUMAN		Т	HOMAN	T	T	ISSEROI	T LI IMANI		
Sir	lar t Top Hit Acess	Value	6.0F-04 AWARDE40 4	8 0F-04	5.0E-04 010341	R OF 04 A MORES	.0C-0+ AW631844.1	5.0E-04 AA548931.1		5.0E-04 AF-248054.1	1		50E-04 A 1200000 2	1	1	5 0F-04 At 048607 2	Т	AF248054.1	AA568513.1		192741482.1	232/40.	4.0E-04 AI720263.1 E	4.0E-04 A1720283 4	AW753358 4	Al 163278 2	Al Odrzod 4	Ogenta Dogen	AF281074 4	AV696624 1	AA676331.1	
. }	-	>	2.26	L				1.42 5.	2 10				5.67		L		1.	1		1.40	1		1.57 4.0	1.57 4.05	L		L			0.84 4.0E	2.75 4.0E-04	
	EQ Expression			2	85			35	<u> </u>			_															-			Ö	5	
-	ORF SEQ ID NO:		8		2 20385			3 23085	25113						27608		07720	3			20410		20606	20607	21218	21823		22348	22850	23041	23900	
-	SEQ (D	_1			1	5 11390	L	7 13286	1 15281	3 15801	16227			.		18220	15284	L	L	Ŀ	10592	02207	86/2	10756	11354	11928	11978	12457	13052	13237	14128	
	Probe SEQ ID NO:		9225	8928	939	5		3367	5361	5998	636		6633	7345	7543	8343	8950	916	8944	386	657	2	3	88	2	Ř3	88 88	2588 2588	3127	3316	4228	

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Top Hit Descriptor	nh10a10.s1 NCI_OGAP_Co1 Homo sapiens cDNA clone IMAGE:951930 3' similar to gb:M21121 T-CELL SPECIFIC RANTES PROTEIN PRECURSOR (HUMAN);	Zn61c08.s1 Stratagene muscle 937209 Homo sapiens cDNA clone IMAGE:562670 3'	601345895F1 NIH_MGC_8 Homo sapiens cDNA clone IMAGE:3678910 5'	yy78b10.s1 Soares_multiple_scierosis_2NbHMSP Homo sapiens cDNA clone IMAGE:279643 3' similar to	contains Alu repetitive element;	yx39e12.r1 Soares melanocyte 2NbHM Homo sapiens cDNA clone IMAGE:264142 5	ov87h03.s1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1644341 3/	Mus musculus neuropilin-2(a17) mRNA, alternetively spliced, complete cds	Homo sapiens SMARCA4 isoform (SMARCA4) gene, complete cds, alternatively spliced	FORMIN (LIMB DEFORMITY PROTEIN)	DKFZp761J221_r1 761 (synonym: hamy2) Homo saplens cDNA clone DKFZp761J221 5'	180 KD SECRETORY PHOSPHOLIPASE AZ RECEPTOR PRECURSOR (PLA2-R)	Human short chain acy/ CoA dehydrogenase gene, exons 1 and 2	th23a02.x1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:2119082 3'	INTERNALIN B PRECURSOR	GLUTAMATE DEHYDROGENASE 2 PRECURSOR (GDH)	Homo sapiens Xq pseudoautosomal region; segment 1/2.	PM0-HT0339-190200-007-912 HT0339 Homo sapiens cDNA	QV3-DT0045-221299-046-d09 DT0045 Homo capiens cDNA	V.carteri gene encoding volvoxopsin	Homo sapiens chromosome 21 segment HS210081	Homo sapiens chromosome 21 segment HS210078	FIBROBLAST GROWTH FACTOR RECEPTOR 3 PRECURSOR (FGFR-3)	zx48d08.r1 Soares_testis_NHT Homo seplens cDNA clone IMAGE:795471 5' similer to gb:M62762 [VACUOLAR ATP SYNTHASE 16 KD PROTECLIPID SUBUNIT (HUMAN);	et24g05.s1 Soares_testis_NHT Hamo sapiens cDNA clone 1391288 3' similar to gb:M36072 60S RIBOSOMAL PROTEIN L7A (HUMAN):	Inc38e04.r1 NCI CGAP Pr2 Homo sapiens cDNA clone IMAGE:1010430 similar to contains L1.12 L1	repetitive element;	Homo sapiens mRNA for KIAA0749 protein, partial cds	DKFZp547L185_r1 547 (synonym: hfbr1) Homo sapiens cDNA clone DKFZp547L185 5	Homo sapiens SCG10 like-protein, helicase-like protein NHL, M68, and ADP-ribosylation factor related protein 1 (ARFRP1) genes, complete cds
Top Hit Database Source	EST_HUMAN	EST_HUMAN	EST_HUMAN		EST HUMAN	EST HUMAN	EST HUMAN	NT	ΤN	SWISSPROT	EST_HUMAN	SWISSPROT	Ν	EST_HUMAN	SWISSPROT	SWISSPROT	N	EST_HUMAN	EST_HUMAN	TN	INT	IN	SWISSPROT	EST HUMAN	EST HIMAN		EST HUMAN	N F	EST_HUMAN	NT
Top Hit Acession No.	AA578331.1	AA086324.1	BE560660.1		N48313.1	N26507.1	AI025699.1	AF022855.1	AF254822.1	Q05860	AL119426.1	P49259	U83991.1	AI399674.1	P25147	P49448	AJ271735.1	BE153778.1	AW937723.1	Y11204.1	AL163281.2	AL163278.2	P22607	AA454055.1	AA781201 1		AA228301.1	AB018292.1	AL134483.1	AF217796.1
Most Similar (Top) Hit BLAST E Value	4.0E-04	4.0E-04	4.0E-04				4.0E-04	4.0E-04	4.0E-04	4.0E-04	3.0E-04	3.0E-04	3.0E-04	3.05-04	3.0E-04	3.0E-04	3.0E-04	3.0E-04	3.0E-04	3.0E-04	3.0E-04	3.0E-04	3.0E-04	3.0E-04	305-04	-	3.0E-04	3.0E-04	3.0E-04	2.0E-04
Expression Signal	2.75	1.28	3.72		6.0	1.62	2.79	1.44	1.62	1.84	3.04	2.22	1.27	1.18	3.37	3.63	1.18	4.5	0.85	1.34	4.82	6.88	3.49	1.42	3.66		2.33	4.32	2.01	1.18
ORF SEQ ID NO:	23901	24121	24681		ł	27040					18942	18977	65302		22988				24456			26166	26875	27780	. JANA		24908	25068		19957
Exan SEQ ID NO:	14128	14332	14893	<u> </u>	_1		17439	17512	19505	19479	10124	10160	10788	11707	13180	13783	13876	14597	14669	14924	15843	16026	16885	17556	17827		19726	19578	19361	10141
Probe SEQ ID NO:	4228	4437	5020		5132	6972	7588	7662	9296	2296	150	188	862	1810	3268	3882	3969	4711	4784	5052	5735	6081	9089	7705	7977		9118	9483	9874	169

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Table 4
Single Exon Probes Expressed in Heart

Probe SEQ ID NO: 1162 1163 1781 1781 1781 1781 1781 1781 1781 178	<u> </u>	g _Θ	Sign.	(Top) Hit BLAST E Value 2.0E-04 2.0E-0	Top Hit Acess No. No. No. Hall AU146707.1 M88624.1 M88624.1 AL163203.2 AF224268.1 AL163203.2 AF224268.1 AR124529.1 BE082317.1 U84374.1 U041029.1 H98265.1 H98265.1 AV664352.1 AV664352.1 AV664362.1 AV664362.1 AV664362.1 AV664362.1 AV664362.1 AV664362.1 AV664362.1 AV664362.1 AV664362.1 AV664362.1 AV664362.1 AV664362.1 AV664362.1 AV664362.1 AV664362.1 AV664362.1 AV664362.1 AV664362.1 AV664362.1	Top Hit Database Source Source Source NT NT NT NT EST HUMAN NT EST HUMAN NT EST HUMAN NT EST HUMAN NT EST HUMAN EST HUMAN EST HUMAN EST HUMAN EST HUMAN EST HUMAN EST HUMAN EST HUMAN EST HUMAN EST HUMAN NT EST HUMAN EST HUMAN EST HUMAN EST HUMAN EST HUMAN EST HUMAN NT EST HUMAN EST HUMA	Top Hit Describtor AU146707 HEMBB1 Home septens cDNA clarie HEMBB1001253 3* Human dystrophin gene Human dystrophin gene Human dystrophin gene Human dystrophin gene Human genetitive element; Home septens chromosome 21 segment HS21C003 Mks muscalus 5* Tanking region of Pitk3 gene AU24B7082831* TCRBV1851,
6638	16698	26709	1.33	2.0E-04	U32444.2 AB026898.1	TN TN	Solanum lycopersicum phytochrome F (PHYF) gene, partial cds Homo saplens DNA, DLEC1 to ORCTL4 gene rigion, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes, complete cds) Homo saplens DNA, DLEC1 to ORCTL4 gene rigion, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes,
6818	16698	26891	1.19	2.0E-04	AB026898.1	ΝΤ	complete cds)

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Table 4

Single Exon Probes Expressed in Heart	Top Hit Descriptor	Live of the state	December 3 Press (FRIT) gene, exon 6	RC3-T10Z34-131099-011-b05 HT0Z54 Homo sapiens cDNA	AVENCETIFITY Sources testis NHT Homo sapiens cDNA clone IMAGE:742964 5'	HV730373 RTF Homo Septens cDNA done HTFAAA01 5	1901111XI NCI_CGAP_Gas4 Homo saplens cDNA clone IMAGE:2140269 3' similar to contains Alu remetitive	element;	926099. St. Stoares melanocyte 2NBHM Homo sapiens cDNA clone IMAGE:2717190 3' 11 11 1 renative elements elements and the sapiens cDNA clone IMAGE:262864 3' similar to contains	RETROVIRUS-RELATED POLYPROTEIN (CONTAINS: REVERSE TRANSCRIPTASE	ENDONUCLEASE)	IU-H-BIQ-8ab-e-09-0-UI.s1 NCI_CGAP_Sub1 Homo sapiens cDNA clone IMAGE:2708825 3	A	Anguina anguina copamine D1A1 receptor (d1A1) gene, complete cds	inspecies succenterassociated nespessifius ORF168 gene, partial cds; and ORF 69, kaposin, v-FLIP, v-cyclin, latent nuclear antigen, ORF K14, v-GPCR, putative phosphoribosyfform/glycinamidine synthase, and LAMP (LAMP) genes, commissioned	Kaposi's sarcoma-associated herpesvirus ORF 68 gene, partial cds; and ORF 69, kaposin. v-FLIP v-cyclin	revent increas amagen; OKF K14, v-GPCR, putative phosphoribosyfformytalyoinemidine synthese, and LAMP (LAMP) genes, complete cds	Equus caballus DNA, chromosome 24g14, microsafallile TKV3a	Homo sapiens membrane-bound aminopeotidase P (XNPFP2) nego company	Homo sapiens membrane-bound aminopeptidasia P (XNPEP2) gans commissions	hv45c08.x1 NCI CGAP Lu24 Home serilens activity along the serious customers.	hv45c08.x1 NCI_CGAP_Lu24 Homo sapilers cPNA_closs NAACE:3176366 3	SPLICEOSOME ASSOCIATED PROTEIN 62 (SAP 62) (SPI ICINI) EACTOR 3.4 ELINING EACTOR	#01f11.x1 NCI_CGAP_Gas4 Homo sepiens cDNA clone IMAGE:2140269 3' similar to contains Alu repetitive	Waller alpha 4 to 1/7	Widouse alpha 1 type-IV collegen mRNA AV847727 OLD US	NOGATIZI GLO DOMO Saplens CUNA clone GLOBBD043'	Homo saplens KIAA0237 gene product (KIAA0237) mRNA
gle Exon Prol	Top Hit Database Source	I-N	TOT LIMAN	TOTAL TOTAL	EST HIMAN	NT TOWNS	100	EST HIMAN	EST HIMAN		Т	Т	FIGURAL PIN	┱	L _N		١				EST_HUMAN				T	T HI IMAN	Т	
Sin	Top Hit Acession	2.0E-04 AF020503 1	BF149303 4	2 0F-04 AA405777 4	2.0E-04 AV730373 1	2.0E-04 AJ243213.1	2 0F-04 Al440282 1	4 AW136740.1	H99646.1	044000	1,7,4				AF148805.1		AF148805.1		AF195953.1	AF195953.1		33.1	062203	A1440282 4		-	T	7662015
	Most Similar (Top) Hit BLAST E Value	2.0E-04	20E-04	20F-04	20E-04	2.0E-04	2 OF -04	2.0E-04	1.0E-04	10 TO 1		1.0F-04	1 0F-04		1.0E-04			1.0E-04	1.05.04		1.0E-04		1.0E-04	1.0E-04 /		1.0E-04/A		1.0E-04
	Expression Signal	2.20	1.46	1.71	6.14	1.88	5.57	2.94	86.0	2.43	387	3.87	2.97		3.24		3.24	1.67	1.29	1.28	1.15	1.15	1.04		1.78	1.08	0.85	1.56
	ORF SEQ ID NO:	27054	27801	ĺ	28360		28762	28866	20516	20819	20855	20856			21374		21376	21594	2002	777	77.411	22412	22969	23376	23654	23675	24043	24671
	Exon SEQ ID NO:			17604	18107	18387	18480	18582	10680	10976	11014	11014	11216		11515		11515	#L/11	12/60	425.25	7707	7707	2	13589	13878	13898	14258	14901
	Probe SEQ ID NO:	6982	7729	7754	8225	8515	8625	8726	750	1059	1098	1098	1310		1610		1610	101	2800	28.5	2000	2007	747	3876	3971	3991	4362	5028

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Top Hit Descriptor	Homo sapiens KIAA0237 gene product (KIAA0237), mRNA	qx62h04.x1 NCI_CGAP_GC4 Homo sepiens cDNA clone IMAGE:2005975 3'	S.cerevisiae chromosome VII reading frame ORF YGL038c	LINE-1.REVERSE TRANSCRIPTASE HOMOLIOG	qv57d10.x1 NCI_CGAP_Ov32 Homo saplens cDNA clone IMAGE:1985683 3'	qv57d10.x1 NCI_CGAP_Ov32 Homo sapiens clONA clone IMAGE:1985683 3'	wf28e08.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2356742.3'	CYSTATIN-RELATED EPIDIDYMAL SPERMATOGENIC PROTEIN PRECURSOR (CYSTATIN 8)	Homo sapiens phospholipid scremblase 1 (PLSCR1), mRNA	LINE-1 REVERSE TRANSCRIPTASE HOMOLIOG	Mause alpha leukocyte interferon gene, complete cds	Homo sapiens mRNA for KIAA1142 protein, partial cds	CM0-CT0404-130700-475-h03 CT0404 Homo caplens cDNA	CM2-NN0010-220300-124-d08 NN0010 Homo sapiens cDNA	ah45c11.s1 Soares_testis_NHT Homo saplens cDNA clone 1292468 3'	w54c11.x1 NCI_CGAP_Co16 Homo saplens cDNA clone IMAGE:2394068 3' similar to contains MER6.t1	MER6 repetitive element ;	PROLYL 4HYDROXYLASE ALPHA-2 SUBUNIT PRECURSOR	Homo saplens gene for cholecystokinin type-A receptor, complete cds	Homo sapiens methyl-CpG binding protein 1 (MBD1) gene, exon 15b	xa34g05.x1 NCi_CGAP_Br18 Homo saplens cDNA clone IMAGE:2569728 3' sImilar to contains L1.t2 L1 repetitive element :	qv23f06.x1 NCI_CGAP_Lym6 Hamo sapiens cDNA clone IMAGE:1982435 3' similar to contains element	MIR repetitive element;	PROLYL 4-HYDROXYLASE ALPHA-2 SUBUNIT PRECURSOR	TO THE COLUMN TO	ngmo septens Mondo gene, partiet cas; end culori, JUARI, cob, cob, cob, cob, cob, cob, cob, cob	Pisum sativum mRNA for beta-1,3 glucanase (gns2 gene)	Pisum sativum mRNA for beta-1,3 glucanase (girs2 gene)	Human platelet-derived growth factor A chain (PDGFA) gene, exons only	HSPD16734 HM3 Homo sapiens cDNA clone s3000003H04	Human haptoglobin and haptoglobin-related protein (HP and HPR) genes, complete cds
Top Hit Database Source	NT	EST_HUMAN	LN L	SWISSPROT	EST_HUMAN	EST_HUMAN	EST_HUMAN	SWISSPROT	LN	SWISSPROT	Ę	NT	EST HUMAN	EST_HUMAN	EST_HUMAN		EST_HUMAN	SWISSPROT	IN	LZ	EST HUMAN		EST_HUMAN	SWISSPROT		L Z	NT	N.	L	EST_HUMAN	NT
Top Hit Acession No.	7662015 NT	Al357156.1	Z72560.1	P08547	AI251980.1	A1251980.1	AI806220.1	088969	10863876 NT	P08547	M28587.1	AB032968.1	BE696769.1	AW893325.1	AA718933.1		AI762209.1	Q60718	D85606.1	AF120982.1	AW073078.1		AI287878.1	Q60716		AF129756.1	AJ251646.1	AJ251646.1	M83575.1	F28172.1	M69197.1
Most Similar (Top) Hit BLAST E Value	1.0E-04	1.0E-04 /	1.0E-04	1.0E-04	1.0E-04	1.0E-04	1.0E-04	1.0E-04	1.0E-04		1.0E-04	1.0E-04	1.0E-04	1.0E-04	9.0E-05		9.0E-05	9.0E-05	9.0E-05	9.0E-05	9.05-05		9.0E-05	9.0E-05		9.0E-05				8.0E-05	8.0E-05
Expression Signal	1.58	66.0	6.0	1.36	12.77	14.47	2.53	1.48	1.75	9	2.17	1.98	1.84	1.65	14	-	8.0	1,47	2.9	2.93	. 28		1.85	4.4		417	1.48	3.53	0.78	1.16	1.87
ORF SEQ ID NO:	24672	24676		25563	56283		27455		27628			28012			20439			25632		27470	28617		28705				20576				28630
Exan SEQ ID NO:	14901	14905	15015	15487	16129	16129	17249	17253	17413	17698	18513	18721	18824	19158	10616		13902	15543	17264	17266	18352		18436	15543		19508	10733	10773	12847	15080	18364
Probe SEQ ID NO:	5028	5033	5148	5572	6264	6468	7380	7384	7562	7848	8649	8913	8035	9298	883		3995	5628	7455	7457	8479		8568	8890		8328	804	848	2920	4852	8491

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Top Hit Descriptor	zs88h01.s1 NCI_CGAP_GCB1 Homo sapiens ¿DNA clone IMAGE:704593 3' similar to contains Alu repetitive element,contains element MSR1 repetitive element;	RC3-CT0208-220999-011-E04 CT0208 Homo sapiens cDNA	RC3-CT0208-220999-011-E04 CT0208 Homo sapiens cDNA	HUM072014F Human fovea cDNA Homo sapieris cDNA clone EST HFD072014	HUM072014F Human fovea cDNA Homo sapieris cDNA clone EST HFD072014	PROBABLE GLYCEROL-3-PHOSPHATE ACYLTRANSFERASE, MITOCHONDRIAL PRECURSOR (GPAT)	Hamo sapiens chromosome 21 segment HS21C078	Dictyostelium discoldeum gene for TRFA, complete cds	Hano sepiens chromosome 21 segment HS21C001	Ceenorhabditis elegans Skp1p homolog mRNA, complete cds	Rat cytomegalovirus Maastricht, complete genome	EST78713 Placenta I Homo sapiens cDNA	EST04984 Fetal brain, Stratagene (cat#936206) Homo sapiens cDNA clone HFBED60	Hamo sapiens sarcoglycan, epsilon (SGCE), mRุNA	Hamo sapiens chromosome X open reading frame 6 (CXORF6) mRNA	Homo saplens chromosome X open reading frame 6 (CXORF6) mRNA	wb54h08.x1 NCI_CGAP_GC6 Homo sapiens cDNA clone IMAGE:2308531 3' similar to gb:J03250 DNA TOPOISOMERASE I (HUMAN);	H. saplens flow-sorted chromosome 6 Hindill fragment, SC6pA28B10	H.saplens flow-sorted chromosome 6 Hindill fragment, SC6pA28B10	Homo sapiens monocyte/neutrophil elastase inhibitor gene, complete cds	CONTACTIN PRECURSOR (GLYCOPROTEIN GP135)	CONTACTIN PRECURSOR (GLYCOPROTEIN GP135)	yv50g11.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:246212 5'	PM4-NN0050-310300-001-f10 NN0050 Homo sapiens cDNA	C4B-BINDING PROTEIN PRECURSOR (C4BP)	C4B-BINDING PROTEIN PRECURSOR (C4BP)	ye28c12.r1 Stratagene lung (#937210) Homo sapiens cDNA clone IMAGE:119062 5'	y59408.s1 Sogres placenta Nb2HP Homo sapiens cDNA clone IMAGE:143535 3' similar to contains Alu	repositive element; contains LTK/ repetitive element;	zk58t02.r1 Soares_pregnant_uterus_NbHPU Homo sapiens cDNA clone IMAGE:487036 5	MR0-N10038-250400-001-109 N10038 Homo sepiens cDNA
Top Hit Database Source	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	SWISSPROT	NT.	LN	TN	IN	LN LN	EST_HUMAN	EST_HUMAN	NT	L	NT	EST HUMAN	N	IN	NT	SWISSPROT	SWISSPROT	EST_HUMAN	EST_HUMAN	SWISSPROT	SWISSPROT	EST_HUMAN		EST HUMAN	EST HOWAN	EST_HUMAN
Top Hit Acession No.	AA279333.1	AW847445.1	AW847445.1	L49075.1	L49075.1	022849	AL163278.2	AB009080.1	AL163201.2	U60980.1	9845300 NT	AA367812.1	T07095.1	10835046 NT	4885170 NT	4885170 NT	AI655241.1	Z84506.1	Z84506.1	AF053830.1	Q12860	Q12860	N72829.1	AW896629.1	P08607	P08607	T94149.1		R75639.1	AA044015.1	AW890110.1
Most Similar (Top) Hit BLAST E Value	8.0E-05	7.0E-05	7.0E-05	7.0E-05		7.0E-05				7.0E-05	7.0E-05	7.0E-05	7.0E-05	7.0E-05	6.0E-05	6.0E-05	6.0E-05	6.0E-05	6.0E-05	6.0E-05	6.0E-05		ľ	6.0E-05	6.0E-05	6.0E-05	6.0E-05		6.0E-05		6.0E-05
Expression Signal	3.45	2.9	2.9	1.05	1.05	2.32	5.24	5.78	1.27	0.82	0.84	1.12	3.04	2.89	1.67	1.57	1.42	0.89	0.89	2.45	3.33	3.33	1.49	2.61	1.37	1.37	1.28		3.68	3.69	8.44
ORF SEQ ID NO:		20118	20119	20302	20303	20800	22440	22843	23953	24023	24508		27505		21766	21767	22309	22409	22410	20415	25599	25600	25879	26989	27437	27438	27558			28911	25073
SEQ ID NO:	19589	10303	10303	10498	10496	10957	12551	13046	14175	14241	14723	16065	17296		11874	11874	12418	12519	ı	li	15519		i	16796		17234	17354		18019	18621	19587
Probe SEQ ID NO:	8911	344	344	555	555	1039	2686	3121	4276	4344	4842	5202	7508	8501	1981	1981	2544	2652	2652	27.83	5605	5605	5855	8918	7330	7330	7484		8131	ò	9534

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Vatue	Top Hit Acession No.	Top Hit Database Source	ာ်စp Hit Descriptor
9965	19427		1.54	6.0E-05	BE858403.1	EST_HUMAN	7g28a08.x1 NCI_CGAP_Bm23 Homo saplens cDNA clone IMAGE:3307766 3'
1382	11287	21141	14.37		AW392086.1	EST_HUMAN	QV4-ST0234-241199-040-h11 ST0234 Homo sapiens cDNA
1819	11716		1.63	5.0E-05	TN 168523891	LN	Homo sapiens 22kDa peroxisomal membrane protein-like (LOC55895), mRNA
2831	12760	22550	0.88	5.0E-05	AJ251058.1	۲N	Homo sapiens MEP1A gene, promoter region and exon 1
3897	13807	23593	2.89	5.0E-05	AJ251884.1	N	Homo sapiens partial SLC22A3 gene for extraneuronal monoamine transporter (EMT), exon 1
5392	15311	25165	9.01	5.0E-05	X58855.1	NT	Human MLC1emb gene for embryonic myosin alkaline light chain, 3'UTR
5646	15559	25652	3.46	6.0E-05	AV653544.1	EST_HUMAN	AV653544 GLC Homo sapiens cDNA clone GLCDMA08 3'
8328	19173		2.96	5.0E-05	P49183	SWISSPROT	RETINAL-BINDING PROTEIN (RALBP)
9590	19173		3.47	5.0E-05	P49183	SWISSPROT	RETINAL-BINDING PROTEIN (RALBP)
2776	10193		4.68		U12821.1	NT.	Human renin (REN) gene, 5' flanking region
7486	17358		71.7	4.0E-05	AF202635.1	F	Homo sapiens PP1200 mRNA, complete cds
							hi36c07.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2974380 3' similar to contains
8150	18038	28286	4.6	4.0E-05	AW627946.1	EST_HUMAN	element MIR repetitive element ;
9287	18989		2.17	4.0E-05	AW117580.1	EST_HUMAN	xd83e09.x1 Soares_NFL_T_GBC_S1 Homo expiens cDNA clone IMAGE:2805192 3'
							qh84c10.x1 Soares_fetal_liver_spleen_1NFLS.US1 Homo sapiens cDNA clone IMAGE:1849458 3' similar to
685	10599	20417	0.84		AI248061.1	EST_HUMAN	contains Alu repetitive element;contains element KER repetitive element ;
1043	10961	20804	1.36	3.0E-05	AW273851.1.	EST_HUMAN	xx24g03.x1 Soares_NFL_T_GBC_S1 Homo sepiens cDNA clone IMAGE:2814100 3'
1115			1.28	3.0E-05		EST_HUMAN	601461463F1 NIH_MGC_66 Homo saplens cDNA clone IMAGE:3865142 5'
1115		20871	1.28	3.0E-05	BF037898.1	EST_HUMAN	601461463F1 NIH_MGC_68 Hamo sapiens cDNA clone IMAGE:3865142 5'
2688	12553	22442	0.91	3.0E-05	Q62234	SWISSPROT	SKELEMIN
4286	14184	23964	68.89	3.0E-05	BE169211.1	EST_HUMAN	PM1-HT0521-120200-001-e10 HT0521 Homo sapiens cDNA
4285	14184	23965	6.89	3.0E-05	BE169211.1	EST_HUMAN	PM1-HT0521-120200-001-e10 HT0521 Hamo saplens cDNA
4365	14261	24045	1.19	3.0E-05	AA368679.1	EST_HUMAN	EST79896 Placenta I Homo saplens cDNA similar to similar to p53-associated protein
4385	14261	24046	1.19	3.0E-05	AA368679.1	EST_HUMAN	EST79996 Placenta I Homo saplens cDNA similar to similar to p53-associated protein
4518	14412	24197	0.85	3.0E-05	AF149773.1	IN	Homo sapiens NOD1 protein (NOD1) gene, excins 1, 2, and 3
							qh64c10.x1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:1849458 3' similar to
4731	10599	20417	0.84	3.0E-05	A1248061.1	EST_HUMAN	contains Alu repetitive element; contains element KER repetitive element;
5151		24786	0.97	3.0E-05	AV726630.1	EST_HUMAN	AV726830 HTC Hamo saplens cDNA clone HTCCEA01 5
6412	16332	25382	1.64	3.0E-05	11072102 NT	NT	Mus musculus myosin light chain 2, precursor lymphocyto-specific (Mylc2pl), mRNA
6811	16491	26677	2.23	3.0E-05	BE733157.1	EST_HUMAN	601567451F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3842292 5'
6863	16742	26935	1.62	3.0E-05	AA284049.1	EST_HUMAN	zs60b05.s1 Stratagene schizo brain S11 Homo sapiens cDNA clone IMAGE:701841 3'
7154			1.65	3.0E-05	AW770982.1	EST_HUMAN	hl94e08.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3009638 3'
7158				3.0E-05	6912431 NT	NT	Homo saplens interleukin-1 receptor antagonist homolog 1 (IL1HY1), mRNA
7373	17242	27447	1.33	3.0E-05	AA372562.1	EST_HUMAN	EST84475 Colon adenocarcinoma IV Homo saplens cDNA 5' end

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Table 4
Single Exon Probes Expressed in Heart

Most Similar (Top) Hit Acession Detabase Value BLAST E No. Source Source Value 3.0E-05 A1769331.1 EST_HUMAN 2.0E-05 M13792.1 NT EST_HUMAN 2.0E-05 M13792.1 NT EST_HUMAN 2.0E-05 M13792.1 NT EST_HUMAN 2.0E-05 A1263349.1 EST_HUMAN 2.0E-05 A1263349.1 EST_HUMAN 2.0E-05 A1263349.1 EST_HUMAN 2.0E-05 A1263349.1 EST_HUMAN 2.0E-05 A1263349.1 EST_HUMAN 2.0E-05 A1263349.1 EST_HUMAN 2.0E-05 A126325.1 EST_HUMAN 2.0E-05 A126325.1 EST_HUMAN 2.0E-05 A126325.1 EST_HUMAN 2.0E-05 A126326.1 NT 2.0E-05 A126301.1 EST_HUMAN 2.0E-05 A126301.1 EST_HUMAN 2.0E-05 M41751.1 EST_HUMAN 2.0E-05 M41751.1 EST_HUMAN 2.0E-05 A1391025.1 EST_HUMAN 2.0E-05 A1391025.1 EST_HUMAN 2.0E-05 A1391025.1 EST_HUMAN 2.0E-05 A1391025.1 EST_HUMAN 2.0E-05 A1391025.1 EST_HUMAN 2.0E-05 BE175801.1 EST_HUMAN 2.0E-05 BE175801.1 EST_HUMAN 2.0E-05 BE175801.1 EST_HUMAN 2.0E-05 BE175801.1 EST_HUMAN 2.0E-05 BE175801.1 EST_HUMAN 2.0E-05 BE175801.1 EST_HUMAN 2.0E-05 BE175801.1 EST_HUMAN 2.0E-05 BE175801.1 EST_HUMAN 2.0E-05 BE175801.1 EST_HUMAN 2.0E-05 BE175801.1 EST_HUMAN 2.0E-05 BE175801.1 EST_HUMAN 2.0E-05 BE175801.1 EST_HUMAN 2.0E-05 BE175801.1 EST_HUMAN 2.0E-05 BE175801.1 EST_HUMAN 2.0E-05 BE175801.1 EST_HUMAN 2.0E-05 BE175801.1 EST_HUMAN 2.0E-05 BE175801.1 EST_HUMAN 2.0E-05 BE175801.1 EST_HUMAN 2.0E-05 BE175801.1 EST_HUMAN 2.0E-05 A1391025.1 EST_HUMAN 2.0E-05 A		1		1			1								, —			_	-		_					_				_
Expn Mcst Smilar Most Smilar Top Hit Acession Top Hit Acession Top Hit Acession 17421 3.16 3.06-05 Ai768331.1 EST_HUMAN 12160 22056 1.76 2.06-05 Ai768331.1 EST_HUMAN 12242 22302 3.02 2.06-05 Ai768331.1 EST_HUMAN 13255 23027 0.94 2.06-05 Ai768331.1 EST_HUMAN 13243 23027 0.94 2.06-05 Ai768331.1 EST_HUMAN 13243 23027 0.94 2.06-05 Ai768331.1 INT 13866 24453 0.94 2.06-05 Ai286021.1 INT 13849 1.14 2.06-05 Ai286021.1 INT 14460 24453 0.81 2.06-05 Ai28607.1 EST_HUMAN 14460 24453 0.81 2.06-05 Ai28471.1 EST_HUMAN 14460 24453 0.81 2.06-05 Ai28334.1 EST_HUMAN 15872 2602 2.06-05 Ai24330.1 EST_HUMAN 15872 2602 2.06-05 Ai24362.1 INT	יים בילקו פספפת זון חפשון	Top Hit Descriptor	WG36f09x1 Soares NSF F8 9W OT PA IS G HOME STATES THE STATES TANKS THE STATES	qh88e11.x1 Soares_NFL_T_GBC_S1 Homolegapiens cDNA clone IMAGE:1855052 3' similar to contains	Human adenosina desminase (ADA) dans complete ade	2448a12.r1 Strategene hNT neuron (#837233) Homo sepiens cDNA clone IMAGE:632734 5' similar to	Colitains Alu repetitive element; contains element L1 repetitive element; R73, BT0340, 400,000,000, L25, BT0340, 400,000,000, L25, BT0340, 100,000,000, L25, BT0340, 100,000,000, L25, BT0340, 100,000,000, L25, BT0340, BT0340	Homo septems 247 abov VIOE43	H septemb DNA for and consultation of the complete cas	Sparakiae 128 Khr frammat 26th Let	DKFZp5661084 71 566 (synonym; hRd2) Homo seniens cDNA class DKEZ_Essino 1 5	4913aD8.x1 Soares, NhHMPu_S1 Homo sapiens cDNA clone IMAGE:1932374.3' similar to contains	MENTIONS MENTION REPORTING COMMENT;	NOTABLE SOUTH IN IN INTERCATE THE MAIN SERVICES CLINA Clone IMAGE: 3608653 5'	activation of the constraints of	Tronio septens TNN 11 gene, exons 1-11 (and joined CDS)	Model E. 31 No. JOSAP JOST Homo Saplens CUNA Clone IMAGE:1238519 31	MISSENT MINISTER DISPLANCE OF MINISTERS	modelion at located Discharge Colon, NHOW Homo sapiens cDNA clone IMAGE:2522077 3'	Heterodombus francisci Hoxa10 (Hoxa40), Hoxa9 (Hoxa9), Hoxa7 (Hoxa7), Hoxa8 (Hoxa8), Hoxa5 Hoxa8) Hoxa8 (Hoxa8) Hoxa8	", "" (""), "	deterodontus francisci Hoxa10 (Hoxa10), HoxA9 (Hoxa9), HoxA7 (HoxA7), HoxA6 (HoxA6), HoxA5	azonos x1 NCI CASAP CI I Home serior and serior serior serior serior serior serior serior serior serior serior	Mode of R Control along to the control of the contr	TOTALON I SCALES, PIRCEITA, OLDSWORKS, ZNDHPStO9W Homo sapiens cDNA clone IMAGE: 259570 5'	w91a06.r1 Soares placenta 8to9weeks 2NbHp8trotW Home carriens and Anna carriers and	1035h07.x1 Soares Dieckaraefe rolan NHCDillomo control separate curva carte invoces 259570 5	CEHT0582-280300-012-E12 HT0582 Home carriers CDNA GIONE IMAGE: 2522077 3	hw21a03.x1 NCI_CGAP_KId11 Home septems cDNA clone INAGE:3183532.3' similar to TR:Q12832 Q12832 GLYCOPHORIN HEP2	
Expn NO: ORF SEQ ID NO: Expression Signal (Top) Hit Top Hit Aossic Signal Most Similar Top Hit Aossic No. Top Hit Aossic No. 17421 3.15 3.0E-05 Al769331.1 12549 1.78 2.0E-05 Al769331.1 12549 1.78 2.0E-05 Al769331.1 12549 7.21 2.0E-05 Al769331.1 13025 2.2821 1.78 2.0E-05 Al769331.1 13026 2.2821 1.78 2.0E-05 Al769331.1 13026 2.2821 1.78 2.0E-05 Al76952.1 13026 2.2821 1.34 2.0E-05 Al76952.1 13266 2.2821 1.34 2.0E-05 Al76952.1 13266 2.4453 0.81 2.0E-05 Al26341.1 14403 24453 0.81 2.0E-05 Al26341.1 15898 2.602 2.19 2.0E-05 Al76310.1 15898 2.602 2.19 2.0E-05 Al7630.1 15898 2.602 2.19 2.0E-05 Al7630.1 16109 2.6260 2.19 2.0E-05 Al7630.1 16409 2.6260	910	Top Hit Database Source	EST HUMAN	EST HIMAN	N	NAMIL TOT	EST HIMAN		Ę	LZ	EST_HUMAN	MAN IJ FAR	TOT LINAMAN	- FZ				T HI MAN	Т				T HUMAN		Т		Г	П		
Expn NO:: ORF SEQ ID NO:: Expression Signal Most Similar (Top) Hit BLAST E 17421 3.15 3.0E-05 12412 22058 1.78 2.0E-05 12546 7.21 2.0E-05 13225 2302 3.02 2.0E-05 13243 23027 0.94 2.0E-05 13243 23050 1.14 2.0E-05 13849 0.85 2.0E-05 14460 1.06 2.0E-05 14866 24453 0.89 2.0E-05 15898 26022 1.9 2.0E-05 15898 26022 2.19 2.0E-05 15983 26260 2.19 2.0E-05 16108 26260 2.19 2.0E-05 16109 26260 2.21 2.0E-05 16109 26261 2.21 2.0E-05 16383 2.8071 2.45 2.0E-05 17894 28141 2.81 2.0E-05 17897 28141 2.98		Top Hit Acession No.	AI769331.1	A1286021 1					X89211.1	X95465.1			T	Ī	T			Ţ										Γ	BE348229.1	
Exan SEQ ID NO: Signa NO: 17421		Most Similar (Top) Hit BLAST E Value	3.0E-05	2.0E-05	2.0E-05				2.0E-05	2.0E-05	2.0E-05	2.0E-05	2.0E-05	2.0E-05	2.0E-05	2.0E-05	2.0E-05	2.0E-05				2.0E-05/	2.0E-05/				2.0E-06	2.0E-05	2.0E-05	
Exan SEQ ID ORI SEQ ID ORI NO: 17421 17421 17421 13243 13366 14480 14668 15488 15972 16483 2 17874 2 15983 17897 2 15983 17897 2 15983 17897 2 15983 17897 2 15983 17897 2 15983 17897 2 15983 17897 2 15983 17897 2 15983 17897 2 15983 17897 2 19514		Expression Signal	3.15	1.78	3.02	7.21	1.6	0.94	1.14	٦	0.85	08.0	1.08	0.81	1.42	2.19	2.19	8.04		2.21		2.24	2.3	2.45	,	2.45	2.42	2.83	2.98	
									23050			24191		24453	25502	28022	26108	-		26260		26261	26671	28116	1 1 1 1	71187		28141		
Probe SEQ ID NO: 7570 7570 7570 2278 2538 3323 3304 3323 3324 3410 4592 4782 6520 6520 6520 6520 6523 6524 6524 6524 6524 6524 6524 6524 6524		Econ SEQ ID NO:	17421	12160	12412	12546	13025	13225	13243	13366	13849	14403	14480	14668	15438	15898	15972	15983		16109		16109	16483	17874	17071	\$6	15983	17897	19514	
		Probe SEQ ID NO:	7570	2276	2538	2681	3088	3304	3323	3449	3737	4510	4592	4782	5520	5933	6125	6136		6243		6243	6803	8024	7608	*700	8039	8748	9335	

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								_							_										
Top Hit Descriptor	xa89e03.x1 NCI_CGAP_Co17 Homo sapiens ::DNA clone IMAGE::2573932 3' similar to contains L1.b3 L1 repetitive element ;	Homo sapiens ABCA1 (ABCA1) gene, complete cds	AU131513 NT2RP3 Homo sapiens cDNA clone NT2RP3002707 6	Homo sepiens chromosome 21 segment HS21,0082	Drosophila melanogaster strain Lamto 120 Suppressor of Hairless (Su(H)) gene, partial cds	MOSAIC PROTEIN LGN	Homo sapiens chromosome 21 segment HS21C003	zw69g04.r1 Soares_testis_NHT Homo saplens cDNA clone IMAGE:781494 5	xy48g11.x1 NCI_CGAP_Lu34.1 Homo sapiens cDNA clone IMAGE:2856548 3'	Homo sapiens chromosome 21 segment HS21C046	Homo saplens Spast gene for spastin protein	ns19g02.s1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:1184114 3' similar to contains L1.t1 L1	L1 repetitive ekament ;	Homo sapiens phospholipase A2, group X (PLA2G10) mRNA, and translated products	52 KD RO PROTEIN (SJOGREN SYNDROME TYPE A ANTIGEN (SS-A)) (RO(SS-A))	Homo sapiens chromosome 21 segment HS210027	zx35h12.s1 Soares_total_fetus_Nb2HF8_9w Homo saplens cDNA clone IMAGE:788519 3' similar to gb.102932 PEROXISOME PROLIFERATOR ACTIVATED RECEPTOR ALPHA (HUMAN);	zs05e11.r1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:684332 5' similar to contains Alu repetitive element contains element TAR1 repetitive element:	UI-H-812-sqk-e-08-0-UI:s1 NCI CGAP Sub4 Homo sapiens cDNA clone IMAGE:2724398 3'	UI-H-BI2-agk-a-08-0-UI:s1 NCI_CGAP_Sub4 Homo sapiens cDNA clone IMAGE:2724398 3'	ha07c10.x1 NOI_CGAP_Kid12 Homo sapiens cDNA clone IMAGE:2873010 3' similar to contains L1.22 L1 repetitive element;	Human hareditary haemochromatosis region, histone 2A-like proten gene, hereditary haemochromatosis (HLA-H) gene, RoRet gene, and sodium phosphiate transporter (NPT3) gene, complete cds	Human hereditary haemochromatosis region, histone 2A-like protein gene, hereditary haemochromatosis (HLA-H) gene, RoRet gene, and sodium phosphate transporter (NPT3) gene, complete cds	tt73a06.x1 NCI_CGAP_HSC3 Homo saplens cONA clone IMAGE:2246386 3'	qg11b08.x1 Soares_placenta_8to9weeks_2NbHP8to9W Homo sapiens cDNA clone IMAGE:1759191 3'
Top Hit Database Source	EST_HUMAN	N.	EST_HUMAN	Į,	FN	SWISSPROT	LN	EST_HUMAN	EST_HUMAN	N ₁	Z		EST_HUMAN	LN	SWISSPROT	TN	EST_HUMAN	FST HIMAN	EST HUMAN	EST HUMAN	EST HUMAN	Į.	۲N	EST_HUMAN	EST_HUMAN
Top Hit Acession No.	AW074604.1	AF275948.1	AU131513.1	AL163282.2	AF088273.1	P81274	AL 163203.2	AA431118.1	AW419134.1	AL163246.2	AJ246003.1) :	AA641846.1	4505844	P19474	4L163227.2	AA452578.1	AA238110 1			AW 466995.1	J91328.1	U91328.1	AI583811.1	A1218983.1
Most Similar (Top) Hit BLAST E Value		2.0E-05/		1.0E-05/							1.0E-05/		-	1.0E-05	1.0E-05	1.0E-05/	1.0E-05	1 05-05	1.0E-05	1.0E-05/			1.0E-05	9.0E-06	9.0E-08
Expression Signal	3.73	2.25	1.81	1.61	1.67	9.24	1.09	2.01	2.16	4.48	1.42		3.54	6.45	1.8	2.18	2.48	11.8	12	1.2	1.78	22	2.2	6.59	3.66
ORF SEQ ID NO:			25257		23301		Ľ			L	26082		١	26228			27312	27420	L	87772		28414	28415	22398	22775
Exan SEQ ID NO:	19677	19503	19208	12729	13513			14080	14627	14763	15952				16430	17045	17117	17220	Į.	l	17701	18170	18170	12504	12984
Probe SEQ ID NO:	9442	9486	9837	2663	3599	3886	4082	4180	4742	4882	6049		6211	6213	6572	7168	7240	7352	7704	7704	7851	8291	8291	2637	3057

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Г		Г	Г	Τ-	_	Г	I_	<u></u>		Г	Т	Г	Г	1	Г	Т		Ð.	T	Т	T	Г							Г			П
	Top Hit Descriptor	Human alanine/glyoxy/ate aminotransferase (AGXT) gene, exons 1 and 2	Homo sapiens differentiation antigen CD20 gene, exons 5, 6	ox20g01.x1 Soares_fetal_iiver_spleen_1NFLSi_S1 Homo sapiens cDNA clone IMAGE:16569123' similar to	contains Alu repetitive element:	Homo sapiens chromosome 21 segment HS21 C009	SUSHI REPEAT-CONTAINING PROTEIN SRPX PRECURSOR (DRS PROTEIN) (DOWN-REGULATED BY V-SRC)	SUSHI REPEAT-CONTAINING PROTEIN SRPX PRECURSOR (DRS PROTEIN) (DOWN-REGULATED	BY V-SRC)	Human apolipoprotein E (APOE) gene, hepatic control region HCR-2	PUTATIVE SERINE/THREONINE-PROTEIN KINASE C22E12.14C	RC3-CT0283-201199-011-h11 CT0283 Homo saplens cDNA	ANKYRIN 1 (ERYTHROCYTE ANKYRIN)	ANKYRIN 1 (ERYTHROCYTE ANKYRIN)	ab90f10.s1 Stratagene lung (#937210) Homo sapiens cDNA clone IMAGE:854251 3' similar to contains	MERZUTT MERZUT EPERING EIEMENT	Homo sapiens KIAAU555 gene product (KIAAU555), mKNA	qw16g09.x1 NCI_CGAP_Ut3 Homo sapiens cDNA clone IMAGE:1991296 3' similar to contains Alu repetitive lelement:	EST99205 Thyroid Homo saplens cDNA 5' end similar to EST containing L1 repeat	QV2-OT0062-250400-173-h01 OT0062 Homo saplens cDNA	601881522F1 NIH_MGC_57 Homo saplens cDNA clone IMAGE:4093972 5'	QV3-BT0379-010300-105-d11 BT0379 Homo saplens cDNA	QV3-BT0379-010300-105-d11 BT0379 Homo (saplens oDNA	OVARIAN ABUNDANT MESSAGE PROTEIN (OAM PROTEIN)	ox08e02.x1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:1655738 3' similar to	contains MER8.t2 MER8 repetitive element;	Mus musculus E-cadherin binding protein E7 mRNA, complete cds	IL5-UM0070-110400-063-g02 UM0070 Homo saplens cDNA	Homo sapiens calcium channel, voltage-dependent, alpha 11 subunit (CACNA11), mRNA	Homo sapiens chromosome 21 segment HS21C046	Human ABL gene, exon 1b and intron 1b, and putative M8604 Met protein (M8604 Met) gene, complete cds	EST185496 Colon carcinoma (HCC) cell line Homo sapiens cDNA 5' end
	Top Hit Database Source	NT	N		EST_HUMAN	N	SWISSPROT		SWISSPROT	Ν	SWISSPROT	EST_HUMAN	SWISSPROT	SWISSPROT	140711111111111111111111111111111111111	ESI HUMAN	LN.	EST HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	SWISSPROT		EST_HUMAN	NT	EST_HUMAN	N	N	TA	EST_HUMAN
	Top Hit Acession No.	M61755.1	L23416.1		A1034370.1	AL163209.2	Q63769		Q63769	U35114.1	Q10364	AW362539.1	002357	Q02357	, 001000	AA669729.1	7662177 NT	AI368252.1	AA385542.1	AW883141.1	BF215972.1	BE069189.1	BE069189.1	Q01456		AI040099.1	AF167441.1	AW801912.1	11418157 NT	AL163246.2	U07561.1	AA313620.1
	Most Similar (Top) Hit BLAST E Value	90E-06	8.0E-08		9.0E-06	9.0E-08	9.0E-06		9.0E-06	9.0E-06	90E-06	8.0E-06	8.0E-06	8.0E-08	00 10 1	7.0E-06	7.0E-06	7.05-06		7.0E-06	7.0E-06	6.0E-06	6.0E-06	6.0E-06		6.0E-06	6.0E-06	6.0E-06	6.0E-06	5.0E-06	5.0E-06	
	Expression Signal	2.94	2.58		10.24	1.16	2.81		2.81	4.58	3.35	1.55	2.24	2.24	,	1.7	3.19	99.9	0.85	5.73	5.39	1.09	1.02	1.77		2.01	1.46	1.88	1.8	3.58	1.98	6.9
	ORF SEQ ID NO:		26591				27257		27258	27394	28434	22251	29087	29088			21191						23336	22637			24979		26195	25706	25843	27807
	Exan SEQ ID NO:	13473	15513		16428	16809	17069	ı	17069	17193	18186	12722	18796	18796	0000,	10886	11326	12772	1	15412	19698	12814	13549	12838		14539	15203	17520	19377	15604	15731	1 1
	Probe SEQ ID NO:	3569	5599		6570	6831	7192		7192	7317	8309	2483	8882	8392	- 8	3	1420	2844	3516	5493	8072	2887	3635	4647		4663	5281	7670	0066	5695	5825	7817

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Probe		ORF SEQ	Expression	Most Similar (Top) Hit	Top Hit Acession	Top Hit	was Uit Consistent
	NO S	Ö NÖ:	Signal	BLAST E Vælue	o Z	Source	i de mit Descriptor
9795	19311	25204	4.74	5.0E-08	A1085045.1	EST_HUMAN	HA0877 Human fetal liver cDNA library Homo sapiens cDNA
i i	l		7 4 44	A 0F 06	D16267 1	FST HIMAN	ya48c03.r1 Soares infant brain 1NIB Homo sapiens cDNA clone IMAGE:53254 5' similar to contains Alu renetitive element contains L1 repetitive element:
3	L						xc69a12.x1 NCI CGAP Eso2 Homo sapiens cDNA clone IMAGE 2589574 3' símilar to contains Alu
828	10755	20605	. 9.3	4.0E-08	AW103354.1	EST_HUMAN	rapetitive element; contains element MER21 repetitive element;
1312	11218		5.18	4.0E-06	Al334928.1	EST_HUMAN	tb33e09.x1 NCI_CGAP_HSC2 Homo saplens cDNA clone IMAGE:2056168 3'
1312	L		5.18	4.0E-06	Al334928.1	EST_HUMAN	tb33e09.xf NCI_CGAP_HSC2 Homo saplens cDNA done IMAGE:2056168 3'
1458	11363		2.23	4.0E-08	BF365612.1	EST_HUMAN	QV2-NT0046-200600-250-h07 NT0046 Homo sapiens cDNA
2218	12104	22008	1.22	4.0E-08	AW015401.1	EST_HUMAN	UI-H-BI0-aat-1-05-0-UI.s1 NCI_CGAP_Sub1 Homo sapiens cDNA clone IMAGE:27104253'
3025	12953	22746	1.39	4.0E-06	AF198349.1	TN	Gallus gallus Dach2 protein (Dach2) mRNA, complete cds
3819	13731	23520	1.78	4.0E-06	AW848295.1	EST_HUMAN	IL3-CT0214-150200-074-B03 CT0214 Homo sapiens cDNA
	l						wi94c10.x1 NCI_CGAP_Brn25 Homo saplens cDNA clone IMAGE:2432562 3' similar to contains element
4702	14588	24378	1.58	4.0E-06	AI886939.1	EST_HUMAN	MERZZ repentive etement;
4840	14721	24504	1.02	4.0E-08	AL163279.2	NT	Homo sapiens chromosome 21 segment HS21/C079
7109	16986		3.1	4.0E-06	AF009660.1	NT	Homo sapiens T cell receptor beta locus, TCRBV7S3A2 to TCRBV12S2 region
8746	17895	28139	3.74	4.0E-08	AB007955.1	TN	Homo sapiens mRNA, chromosome 1 specific transcript KIAA0486
							z34b08.s1 Soares_fetal_liver_spleen_1NFLS_S1 Homo saplens oDNA clone IMAGE:432663 3' similar to
2117	12006	21904	1.27	3.0E-06	AA700562.1	EST_HUMAN	contains L1.11 L1 repetitive element;
							zi34b08.s1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:432663 3' similar to
2117	12006	21905	1.27	3.0E-06	AA700562.1	EST HUMAN	contains L1.t1 L1 repetitive element;
2220	12105		1.37	3.0E-06	AF202635.1	NT	Homo sapiens PP1200 mRNA, complete cds
							ak48g11.s1 Soares_testis_NHT Homo sapiens cONA clone IMAGE:1409252 3' similar to contains LTR1.t3
2891	12818	22610	0.95	3.0E-06	AA868218.1	EST HUMAN	LTR1 repetitive element;
							wizza05.x1 NCI_CGAP_Ut1 Homo sapiens cDNA clone IMAGE:2425616 3' sImilar to TR:060734 060734
3228	13152]	2.14	3.0E-06	AI857779.1	EST_HUMAN	LINE-1 LIKE PROTEIN ; contains L1.tz L1 repetitive element;
3716	13628	23412	1.28	3.0E-08	BE047094.1	EST HUMAN	hq64d12.x1 NCL_CGAP_HN13 Homo sapiens cDNA clone IMAGE:3124151 3
3716	13628	23413	1.28	3.0E-06	BE047094.1	EST_HUMAN	hq64d12.x1 NCI_CGAP_HN13 Hamo saplens cDNA clane IMAGE:3124151 3'
							yb78b10.r1 Stratagene ovary (#937217) Homo sapiens cDNA clone IMAGE:77275 5' similar to contains L1
4376	14272	24063	0.0	3.0E-06	T50266.1	EST_HUMAN	repetitive element
	ŀ						Homo sapiens gene for alphe-1-miorogiobulin-bikunin, exons 1-5 (encoding alpha-1-microglobulin, N-
4461		24146	4.31	3.0E-08		N	terminus.)
6290	16154		1.92	3.0E-06	P08548	SWISSPROT	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG
9494	19110		4.17	3.0E-06	AW385262.1	EST_HUMAN	RC0-LT0001-281199-011-A03 LT0001 Homo caplens cDNA
196	10167		3.28	2.0E-06	P54386	SWISSPROT	HOMEOBOX PROTEIN GOOSECOID

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		5	7	Ţ	Т	Т	7	T,		7		7:	<u> </u>			٦		7		Τ	7	Т	Т	တ	Γ	Τ	T	Ţ	Τ	7
Top Hit Descriptor	POL POLYPROTEIN (CONTAINS: PROTEASE; REVERSE TRANSCRIPTASE; ENDONUCLEASE)	wa04e03.x1 NCI_CGAP_Kid11 Homo sapiens cDNA done IMAGE:2297068 3' similar to contains MER30.01 MER30 repetitive element;	HISTIDINE-RICH GLYCOPROTEIN PRECURSOR	KNOB-ASSOCIATED HISTIDINE-RICH PROTEIN PRECURSOR (KAHRP)	AV657555 GLC Homo sapiens cDNA clone GLCFDB05 3'	pp02e05.r1 Stratagene ovarlan cancer (#937219) Homo saplens cDNA clone IMAGE: 393232 3	Mus musculus gene for odorant receptor A16, complete cds	wjs0b04.x1 NCI_CGAP_Lym12 Hamo sapiens cDNA clone IMAGE:2410063 3	yu37c04.r1 Sceres ovary tumor NbHOT Homo;saplens cDNA clone IMAGE:2339/4 5 similar to g52X/4929 KERATIN, TYPE II CYTOSKELETAL 8 (HUMAN);	PROTEIN MOV-10	hs92f02.x1 NCI_CGAP_Kld13 Homo sapiens cDNA clone IMAGE:3144699 3' similar to contains L1.t2 L1	repetitive element;	ORGANIC CATION/CARNITINE TRANSPORTER 2 (SOLUTE CARRIER FAMILY 22, MEMBER 5) (FIGH) AFFINITY SODIUM-DEPENDENT CARNITINE COTRANSPORTER)	Mus musculus D6MM5E protein (D6Mm5e) mRNA, complete cds	MEROZOITE SURFACE PROTEIN CMZ-8	Homo sapiens chromosome 21 segment HS21C078	206a12.s1 Soares_fetal_liver_splean_1NFLS_S1 Homo sapiens cDNA clone IMAGE:429982 3' similar to	contains Alu repetitive element;	ZIOSa12.s1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:429982 3' similar to	contains Alu repetitive element;	DNA-DIRECTED RNA POLYMERASE III LARGEST SUBUNIT	Homo sapiens p47-phox (NCF1) gene, complete cds	Homo sapiens p47-phox (NCF1) gene, complete cds	Hilman ARI name even 1b and intron 1b, and butative M8604 Met protein (M8604 Met) gene, complete cds	Home engine chromosome 21 segment HS24 C086	POST POST POST POST POST POST POST POST	Homo saplens chromosome 21 segment H321CU63	MR1-BT0800-030700-002-c06 B10800 Homo sapiens culvA	FIBRINOGEN ALPHA/ALPHA-E CHAIN PRECURSOR	qp54e02.x1 NCI_CGAP_Cos Homo sapiens gunha gione innace. Texcorz o
Top Hit Database Source	SWISSPROT	EST HUMAN	SWISSPROT	SWISSPROT	EST_HUMAN	EST_HUMAN	L	EST_HUMAN	EST_HUMAN	SWISSPROT		EST HUMAN	SWISSPROT	LN.	SWISSPROT	IN.		EST_HUMAN	_	EST_HUMAN	SWISSPROT	NT	NT	ţ		2	N	EST HUMAN	SWISSPROT	EST_HUMAN
Top Hit Acession No.	21414	A1672138.1	2.0E-06 P04929	906719	2.0E-06 AV657555.1	2.0E-08 AA173518.1	2.0E-06 AB030898.1	AI819424.1	H62051.1	P23249		BE328232.1	076082	A FOR 43 84 1	P09125			AA034141.1		AA034141.1	P27625	AF184614.1	AF184614.1				AL163285.2	BF333015.1	_	AI347010.1
Most Similar (Top) Hit BLAST E Value	2.0E-06 P21414	2.0E-06	2.0E-06	2.0E-06 P06719			2.0E-08	2.0E-06 A	2.0E-06	2.0E-06 P23249		2.0E-06	1 0 1 -08					1.0E-08		1.0E-06	1.0E-06		1.0E-06		00-20.1		1.0E-06			1.0E-08
Expression Signal	6.27	2.95	214	2.69	1.11	1.5	1.5	90.9	1.65	1.35		2.48	10	ľ				0.93		0.93	1.2	1	4.74			1.38	1.38	4.69	6.11	1.16
ORF SEQ ID NO:		22108			L	23398			27208		L		10843	L	21206	L		21317		21318		21729				24688	24689		26139	
SEO ID NO:	11455	10001	12295	12383	13391	13614	13623	15784	17013			19148				1	L	11459	L	11459	١.	L	1	<u> </u>	l	14915	14915	15157	16001	16692
Probe SEQ ID NO:	1550	32.26	2418	2519	3475	3700	3710	5878	7136	P403		9548		2 2	1435	1507	3	1554		1554	1585	1949	1949		4273	5043	5043	6233	6107	6813

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Table 4
Single Exon Probes Expressed in Heart

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ביינות באניין המספר באלו פספר ביינות ה	Top Hit Descriptor	qv23f06.x1 NCI_CGAP_Lym6 Homo sapiens cDNA clone IMAGE:1982435 3' similar to contains element MIR repetitive element ;	Homo sapiens shox gene, alternatively spliced products, complete cds	Homo sapiens shox gene, alternatively spliced products, complete cds	zo17e08.r1 Stratagene colon (#937204) Homo sapiens cDNA clone IMAGE:587174 5'	204d11.s1 Soares, total fetus, Nb2HF8 9w Homo sapiens cDNA clone IMAGE:785493 3' similar to gb:D26129 RIBONUCLEASE PANCREATIC PRECURSOR (HUMAN);	Homo sapiens chromosome 21 segment HS21,0003	RC4-NT0054-120500-012-b03 NT0054 Homo sapiens cDNA	Homo sapiens ADP/ATP carrier protein (ANT-2) gene, complete cds	Homo sepiens p47-phox (NCF1) gene, complete cds	Homo saplens p47-phox (NCF1) gene, complete cds	Homo sapiens glypican 3 (GPC3) gene, partial cds and flanking repeat regions	Homo capiens glypican 3 (GPC3) gene, partial cds and flanking repeat regions	Homo sapiens chromosome 21 segment HS210081	q82g07.x1 Soares_NhHMPu_S1 Homo sapien; cDNA clone IMAGE:1878876 3'	ql82g07.x1 Soares_NhHMPu_S1 Hamo sapienis cDNA clone IMAGE:1878876 3'	POL POLYPROTEIN (CONTAINS: PROTEASE; REVERSE TRANSCRIPTASE; ENDONUCLEASE)	Homo sapiens UDP-glucuronosyfransferase gene, complete cds	EST05660 Fetal brain, Stratagene (cat#936206) Homo sapiens cDNA clone HFBEN89	Hamo saplens chromosome 21 segment HS21¢080	Homo sapiens membrane interleukin 1 receptor accessory protein (IL1RAP) gene, exons 10 and 11	CM3-CT0277-221099-024-e11 CT0277 Homo saplens cDNA	Homo saplens HLA class III region containing transcin X (tenascin-X) gene, partial cds; cytochrome P450 21- hydroxylase (CYP21B), complement component C4 (C4B) G11, helicase (SKI2W), RD, complement factor B	(8f), and complement component C2 (C2) genés,>	HYPOTHETICAL 24.1 KD PROTEIN IN LEF4-P33 INTERGENIC REGION	7g94f07x1 NCI_CGAP_Co16 Homo sapiens cDNA clone IMAGE:3314149 3' similar to TR:O75920 O75920	4F5L.;	CM4-NN1029-250300-121-h12 NN1029 Homo sapiens cDNA	wh84f10.x1 NCI_CGAP_Kld11 Homo sapiens cDNA clone IMAGE:2385547 3'	EST93615 Supt cells Homo sapiens cDNA 5' end	
מון ווווסעד פוה	Top Hit Database Source	EST_HUMAN	<u>F</u> Z	NT	EST_HUMAN	EST HUMAN	Z	EST_HUMAN	N	TN	IN	TN	. NT	TN	EST_HUMAN	EST_HUMAN	SWISSPROT	NT	EST_HUMAN	TN	뉟	EST_HUMAN		뉟	SWISSPROT		EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	100
5	Top Hit Acession No.	AI287878.1	U82668.1	U82668.1	AA132611.1	AA449257.1	1.0E-06 AL163203.2	AW890841.1	L78810.1	AF184614.1	AF184614.1	AF003529.1	AF003529.1	AL163281.2	AI288596.1	AI288596.1	P21414	AF135416.1	T07770.1	AL163280.2	AF167341.1	AW855558.1		AF019413.1	P41479		BF001867.1	AW903222.1	AI831893.1	AA380630.1	, 000, 001
	Most Similar (Top) Hit BLAST E Value	1.0E-06	1.0E-08	1.0E-06	1.0E-06	1.0E-06	1.0E-06		1.0E-06		1.0E-06	9.0E-07	9.0E-07	9.0E-07	8.0E-07	8.0E-07	8.0E-07	8.0E-07		8.0E-07	7.0E-07	6.0E-07		6.0E-07	6.0E-07		6.0E-07	8.0E-07	5.0E-07	5.0E-07	20.2
	Expression Signal	1.53	3.72	3.72	4.86	3.73	1.52	4.81	2.99	1.84	1.84	1.49	1.49	238	4.49	4.49	7.45	7.89	7.2	4.15	0.94	2.38		241	1.65		1.35	1.81	3.55	2.9	1000
	ORF SEQ ID NO:	27013		27648	27676				25280	21729	21730	20134	20135	28708	24336	24337					21599	21641		22217			27361			 	
	Exon SEQ ID NO:	16822	17433	17433	17460	17488		18720	19077	11844	11844	10314	10314		14547	14547	15504	16547	18703	18838	11719	11766		12319	13802		17156	19853		10959	L
	Probe SEQ ID NO:	6944	7582	7682	6092	7697	7989	8912	9440	9236	9536	357	298	8572	4661	4661	5589	2999	8893	3025	1822	1870		2442	3891		7279	9303	8Z3	1041	2000

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repetitive element contains element 43R repetitive element;
tg06b05.x1 NCI_CGAP_CL1 Homo saplent cDNA clone IMAGE:2107953 3' similar to contains Alurepetitive element;
repetitive element contains element A3R repetitive element;
xx31a02.x1 NCI_CGAP_B118 Homo saplens; cDNA clone IMAGE:2568362 3' similar to gb:X15341
CYTOCHROME C OXIDASE POLYPEPTIDE VIA-LIVER (HUMAN);
CM-B1178-220499-014 B1178 Homo saplens; cDNA
COLLAGEN ALPHA 1(I) CHAIN PRECURSOR tg06b05.x1 NCI_CGAP_CLL1 Homo saptens cDNA clone IMAGE:2107953 3' similar to contains Alu Homo eaplens NOD1 protein (NOD1) gene, exons 4 through 14 and complete cds Top Hit Descriptor Single Exon Probes Expressed in Heart Top Hit Database Source EST_HUMAN EST_HUMAN Top Hit Acession 5.0E-07 AF149774.1 5.0E-07 AI393981.1 5.0E-07 Al393981.1 Most Similar (Top) Hit BLAST E Value 1.7 Expression Signel ORF SEQ ID NO: 26098 24223 26097

14440 15963

4547 6203 6203 6348 7832 8805 8863

SEQ ID

Probe SEQ ID

15963 16211

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XB318UZXI NCI CGAP Br18 Homo sapiens cDNA clone IMAGE:2568362 3' cimiler to ch: 145244	CYTOCHROME C OXIDASE POLYPEPTIDE VIALIVER (HUMAN);	CW-5 11/8-220499-014 BT178 Homo sapiens cDNA	COLLAGEN ALPHA 1(I) CHAIN PRECURSOR	Homo saplens Xq pseudoautosomal reglon; seament 1/2	QV0-CT0383-210400-204-b12 CT0383 Homp sepiens CDNA	ws84h05.x1 NCI_CGAP_Co3 Homo sapiens cDNA clone IMAGE 25nAso7.2	Xy48g11.x1 NCI_CGAP_Lu34.1 Homo saplens cDNA clone IMAGE-2858548.3	wi81b08.x1 NCI_CGAP_Kid12 Homo sapiensicDNA clone IMAGE วิวัตอวักราว	wi81b08.x1 NCI_CGAP_Kid12 Homo sapiens cDNA clone IMAGE 2389703.31	PM1-BN0083-030300-003-e12 BN0083 Homo sapiens cDNA	Human microfibril-associated glycoprotein (MFAP2) gene, putative promoter region and others.	Untranslated expire and institution spliced	Homo sapiens Xq pseudoautosomal region; segment 1/2	Human polymorphic microsatellite DNA	Human igk subgroup I germline gene, exons 1 and 2 N society 200 miles	nt56b09.s1 NCI_CGAP_Ovz Homo sapiens cCNA clone IMACE: Ganes 5	element, contains L1.13 L1 repetitive element;	Human polymorphic microsatellite DNA	MRO-BN0115-020300-001-f11 BN0115 Homo caplens cDNA	MRO-BN0115-020300-001-f11 BN0115 Homo sapiens cDNA	yd50f12.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone M44.GE-144.ege er	HYPOTHETICAL 63.8 KD PROTEIN IN GUT1:RIM1 INTERGENIC REGION PRECLIDEDE	AV650201 GLC Homo sapiens cDNA clone GLCCCD01 3'	we86b12x1 Soares_NFL_T_GBC_S1 Home sapiens cDNA clone IMAGE 2347967 3	yo14h09.s1 Stratagene lung (#937210) Homo saplens cDNA clone IMAGE:80705 3' similar to similar to	yol 4h09.s1 Strategene lung (#837210) Homo conjens characters.	9b:M62392 ARACHIDONATE 12-LIPOXYGENASE (HUMAN)	1.11.2.11
For	EST CHIMAN	EST DOMAIN	SWISSPROT	N	EST_HUMAN	EST HUMAN	EST_HUMAN	EST HUMAN	EST HUMAN	LO HOMAN	H	2	2 !	Z	L		EST HUMAN	- 2	ES HUMAN	EST HUMAN	- 1	- 1	Т	ES! HOMAN	EST HUMAN	\top	EST HUMAN	
AWOZOBRS 4			A 127479E 4			AW 009602.1	_	_			1119719 1	_	`-	٠_ـــ	M04857.1	A A 670700 4	7,007,00	REODEO774		1047044		V650204 4	707724 4	191500.1	67850.1		5/850.1	
5.0E-07	5.0E-07	5 0F-07	L			4.00-07	4 05 07	4 05-07	4.0F-07		3.05-07	3.0F-07	3 OF 07	20E-07	3.0E-07	2 OF 07	3 0F.07	3 0F-07	3 05-07	3 05 07 7	3 OF 07 Daggar	3.0E-07	3 0F-07 A		3.0E-07 T	100	9.05-0/	
15.83	4.11	4.31	2.08	202	1 08	529	3.84	3.84	2.05		5.01	2.38	2.19	2 23	2	101	1.18	17.25	17.25	0.85	1 82	7,05	0.89		1.47	47.7		
26373	28021	28909			23608	27305	28432	28433			20200	20314	21114				22028	22194	22195	22718	22844	24302	24329	-	24634	24635		
▕	17782	18619	18675	19558	13827	17112	18185	18185	18422		10379	10507	11258	11513		11893	12126	12297	12297	12926	13047	14512	14540	-	14871	14871		
6348	7832	8805	8863	9712	3918	7235	8308	8308	8552		\$	88	1352	1608		2000	2242	2420	2420	2898	3122	4624	4654		4898	4936	1	

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	Top Hit Descriptor	PROTEIN-ARGININE DEIMINASE TYPE IV (PEPTIDYLARGININE DEIMINASE IV) (PAD-R4) (PEPTIDYLARGININE DEIMINASE TYPE AUPHA)	oc04c10.s1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:1339890 3'	QV1-UM0036-200300-115-g02 UM0036 Homp saplens cDNA	Homo saplens chromosome 9 duplication of the T cell receptor beta locus and trypsinogen gene families	Rattus norvegicus mRNA for 45 kDa secretory protein, partial	Homo saplens TRF2-interacting telomeric RAP1 protein (RAP1) mRNA, complete cds	Homo sapiens DiGeorge syndrome critical region, telomeric end	Homo sapiens DiGeorge syndrome critical region, telomento end	Fugu rubripes beta-cytoplasmic(vascular) actin gene, complete cds	Homo saplens homeobox protein CDX4 (CDX4) gene, complete cds and flanking repeat regions	Homo sapiens homeobox protein CDX4 (CDX4) gene, complete cds and flanking repeat regions	208b07.s1 Stratagene NT2 neuronal precurso; 837230 Homo saplens cDNA clone IMAGE:650869 3' similar to gb:L31860 GLYCOPHORIN A PRECURSOR (HUMAN);contains Alu repetitive element;	yc15g04.s1 Stratagene lung (#937210) Homo napiens oDNA clone IMAGE:80790 3' similar to contains L1	יווים ובויא ביווים וויים וויים וויים וויים וויים וויים וויים וויים וויים וויים וויים וויים וויים וויים וויים ו	I/6 AU IOAN IIGEN IIVOATUETIAAI 72 EVA BBOTTEIN CAEZ 48 IN CUBONIOSONIE I	TITOTHER OF THE TOTAL OF THE TO	Homo sapiens caveolin 1 (CAV1) gene, exon 3 and partial cds	xe05h07.x1 Sceres_NFL_T_GBC_S1 Homo sciplens cDNA clone IMAGE:2567485 3' slimitar to WP:C38H2.1 CE00923 PROBABLE RABGAP DOMAINS ;	xa05h07.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2567485 3' similar to WP:C38H2.1 CE00923 PROBABLE RABGAP DOMAINS ;	Homo saplens chromosomie 21 segment HS21C101	RC3-NN0066-260400-021-g11 NN0066 Homo sapiens cDNA	qg58d05.x1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1839177 3'	AV729390 HTC Homo sapiens cDNA clone HTCAEG02 5'	Homo sapiens chromosome 21 segment HS21C103	CM4-NN0003-280300-124-e08 NN0003 Homo saplens cDNA	zn85h11.x5 Stratagene lung carcinoma 937218 Homo sapiens cDNA clone IMAGE:565029 3' similar to contains THR.b2 THR repetitive element ;	Home saplens chromosome 21 segment HS210082
01 1 100	Top Hit Database Source	SWISSPROT	EST_HUMAN	EST_HUMAN	ΙΝ	N	TN	NT	LN	LN	TN	LN T	EST_HUMAN	1	NAMOR - 193	SWISSPROT	SWISSERGI	-Z	EST_HUMAN	EST HUMAN	Į.	EST_HUMAN	EST_HUMAN	EST_HUMAN	LN T	EST_HUMAN	EST_HUMAN	NT
5	Top Hit Acession No.	088807	AA815175.1	AW797168.1	AF029308.1	AJ132362.1	AF262988.1	L77569.1	L77569.1	U38849.1	AF003530.1	AF003530.1	AA223260.1	, 0,000	105042.1	026768	LOSACO	AF125348.1	AW070995.1	AW070995.1	AL163301.2	AW898066.1	A1208715.1	AV729390.1	AL163303.2	AW892507.1	AI732462.1	AL163282.2
	Most Similar (Top) Hit BLAST E Value	3.0E-07	3.0E-07	3.0E-07	3.0E-07	3.0E-07	2.0E-07			. 2.0E-07	2.0E-07	2.0E-07	2.0E-07	_		2.0E-07	2.05-07	2.0E-07	2.0E-07	2.0E-07	2.0E-07	2.0E-07	2.0E-07	2.0E-07	2.0E-07	2.0E-07		1.0E-07
	Expression Signal	10.28	5.32	3	8	2.78	2.62	4.75	4.76	116.1	1.71	1.71	3.11		87.5	1.28	8	14.74	0.97	0.97	0.95	1.73	1.69	3.91	1.61	6.48	1.48	1.21
	ORF SEQ ID NO:	25462		26456			19807	19940			20495	20496	20701			20903	1	23334	24723	24724						77984		
	Exen SEQ ID NO:	15396	15930	16294	18726	19408	10013		10122		10663	10663	10853		L	11060		13547	14948	14948	L	15199		16812	17478	17771	19557	11002
	Probe SEQ ID NO:	5478	9209	6433	8918	666	58	148	148	175	731	731	928	3	יייי אינייייייייייייייייייייייייייייייי	1147	7001	3833	5078	5078	5198	5277	6009	6934	7827	7891	9094	1086

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Probo SEQ ID NO: Expression NO: (Top) Hit Value Signal (Top) Hit Value								
2331 12212 22110 0.94 1.0E-07 P10263 SWISSPROT 2403 1.2280 22177 0.94 1.0E-07 P09266 SWISSPROT 2707 11402 2137 1.0E-07 P09266 SWISSPROT 4185 14095 23374 2.37 1.0E-07 AV718692.1 EST_HUMAN 4185 14095 23875 2.37 1.0E-07 AV718692.1 EST_HUMAN 6103 15897 28475 2.37 1.0E-07 BE047871.1 EST_HUMAN 6424 16285 28473 5.2 1.0E-07 BE047871.1 EST_HUMAN 6478 16285 2847 5.2 1.0E-07 BE047871.1 EST_HUMAN 6778 16861 27786 2.81 1.0E-07 BE047871.1 EST_HUMAN 6782 16861 27780 2.37 1.0E-07 BE047871.1 EST_HUMAN 6782 16861 27780 2.28 1.0E-07 MA1483282.2 NT	Probe SEQ ID NO:		ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
2403 12280 22177 0.94 1.0E-07 7549818 NT 2767 11410 21269 1.76 1.0E-07 AL163282.2 NT 3884 11002 23874 2.37 1.0E-07 AL163282.2 NT 4195 14095 23874 2.37 1.0E-07 AV718682.1 EST_HUMAN 6103 15997 28132 5.2 1.0E-07 AV718682.1 EST_HUMAN 6103 15997 28132 5.2 1.0E-07 BE047871.1 EST_HUMAN 6103 15997 2847 9.04 1.0E-07 BF047871.1 EST_HUMAN 6104 1766 2770 BE-07 BF04785 SWISSPROT 6782 16861 2850 2.81 1.0E-07 BF0435 SWISSPROT 6782 16861 27746 2.77 1.0E-07 BF04435 SWISSPROT 7714 17664 277246 2.81 1.0E-07 BF04455 SWISSPROT 7714	2331	Ц		0.94	1.0E-07	P10263	SWISSPROT	RETROVIRUS-RELATED GAG POLYPROTEIN (VERSION 1)
2797 11410 21269 1.76 1.0E-07 P09266 SWISSPROT 3884 11002 237 1.0E-07 AV718682.1 SST HUMAN 4185 14095 23874 2.37 1.0E-07 AV718682.1 EST HUMAN 6103 15997 28133 5.2 1.0E-07 BE047871.1 EST HUMAN 6103 15997 28133 5.2 1.0E-07 BE047871.1 EST HUMAN 6103 15997 28133 5.2 1.0E-07 BE047871.1 EST HUMAN 6103 15997 28447 9.04 1.0E-07 BE047871.1 EST HUMAN 6782 16861 28851 2.81 1.0E-07 BE04856.1 EST HUMAN 6782 16861 27780 2.37 1.0E-07 BE04856.1 EST HUMAN 7714 17664 27780 2.37 1.0E-07 BE04856.1 EST HUMAN 7716 17680 27780 2.37 1.0E-07 AV734819.1 EST HUMAN 7716 17680 27780 2.81 1.0E-07 AV774450.1 IST HUMAN	2403			0.94	1.0E-07	7549818	IN.	Homo sapiens RAB, member of RAS oncogene family-like 2A (RABL2A), transcript variant 2, mRNA
3684 11002 1.33 1.0E-07 AL183282.2 NT 4185 14095 23874 2.37 1.0E-07 AV718682.1 EST_HUMAN 6103 14096 23875 2.37 1.0E-07 BE04781.1 EST_HUMAN 6103 15997 26133 5.2 1.0E-07 BE04781.1 EST_HUMAN 6103 15997 26133 5.2 1.0E-07 BE04781.1 EST_HUMAN 6782 16861 26850 2.81 1.0E-07 P97435 SWISSPROT 6782 16861 26851 2.81 1.0E-07 P97435 SWISSPROT 6782 16861 26851 2.81 1.0E-07 P97435 SWISSPROT 6782 16861 27246 3.24 1.0E-07 P97435 SWISSPROT 7716 17564 27790 2.37 1.0E-07 P97435 SWISSPROT 7716 17650 27782 1.28 1.0E-07 P97435 SWISSPROT 7716 17650 27782 1.28 1.0E-07 P9746271 EST_HUMAN 8520 <t< td=""><td>2797</td><td></td><td></td><td>1.75</td><td>1.0E-07</td><td></td><td>SWISSPROT</td><td>GLYCOPROTEIN GPV</td></t<>	2797			1.75	1.0E-07		SWISSPROT	GLYCOPROTEIN GPV
4195 14095 23974 2.37 1.0E-07 AV718662.1 EST_HUMAN 4196 14096 23875 2.37 1.0E-07 AV718662.1 EST_HUMAN 6103 15997 28132 5.2 1.0E-07 BE047871.1 EST_HUMAN 6104 16297 28133 5.2 1.0E-07 BE047871.1 EST_HUMAN 6724 162861 28850 2.81 1.0E-07 P97435 SWISSPROT 6782 16881 28851 2.81 1.0E-07 P97435 SWISSPROT 6782 16881 28851 2.81 1.0E-07 P97435 SWISSPROT 6782 16881 28851 2.81 1.0E-07 P97455 SWISSPROT 6782 16881 27782 2.37 1.0E-07 P97455 SWISSPROT 7718 17564 27792 2.37 1.0E-07 P9745541 EST_HUMAN 7708 17830 2.7782 1.56 1.0E-07 P9745241 EST_HUMAN 8520 1783 2.56 1.0E-07 P9746541 EST_HUMAN 8521	3684			1.33			NT	Homo sapiens chromosome 21 segment HS21C082
4195 14096 23875 2.37 1.0E-07 AV718682.1 EST_HUMAN 6103 15697 26132 5.2 1.0E-07 BE047871.1 EST_HUMAN 6103 15697 26133 5.2 1.0E-07 BE047871.1 EST_HUMAN 6424 16285 26847 9.04 1.0E-07 P6681.1 EST_HUMAN 6782 16681 26847 2.01 1.0E-07 P6681.1 EST_HUMAN 6782 16681 27246 3.24 1.0E-07 P6681.1 EST_HUMAN 6782 16681 27726 2.37 1.0E-07 P6681.1 EST_HUMAN 7714 17664 27792 1.28 1.0E-07 A6483678.1 EST_HUMAN 7714 17664 27792 1.26 1.0E-07 A6483678.1 EST_HUMAN 7714 17664 27792 1.56 1.0E-07 A648671.1 NT 9601 17830 2.81 1.0E-07 A648671.1 NT <	4185				1.0E-07		EST_HUMAN	AV718862 GLC Homo saplens cDNA clone GILCFNF04 5'
6103 15897 28132 5.2 1.0E-07 BE047871.1 EST_HUMAN 6103 15897 28133 5.2 1.0E-07 BE047871.1 EST_HUMAN 6103 16897 2813 5.2 1.0E-07 M6081.1 EST_HUMAN 6782 16861 28851 2.81 1.0E-07 P97435 SWISSPROT 7714 17664 277246 2.37 1.0E-07 P67435 SWISSPROT 7714 17669 27724 2.37 1.0E-07 P67435 SWISSPROT 7714 17664 27792 2.37 1.0E-07 P67435 SWISSPROT 7714 17664 27792 2.37 1.0E-07 P67324.1 EST_HUMAN 7714 17669 17780 3.66 1.0E-07 AL163282.2 NT 9497 19113 1.37 1.0E-07 AL163282.2 NT 9661 19222 4.61 1.0E-07 AL163282.2 NT 16851 1933	4195				1.0E-07		EST_HUMAN	AV718682 GLC Homo sapiens cDNA clone GLCFNF04 5'
6103 15997 26133 5.2 1.0E-07 BE047871.1 EST_HUMAN 6424 16285 26447 9.04 1.0E-07 N65081.1 EST_HUMAN 6782 16861 26850 2.81 1.0E-07 P97435 SWISSPROT 77180 17057 27246 3.24 1.0E-07 P67435 SWISSPROT 77181 17054 27726 2.37 1.0E-07 P67435 SWISSPROT 7718 17054 27726 2.37 1.0E-07 P683676.1 EST_HUMAN 7718 17589 27792 1.58 1.0E-07 AL163282.2 NT 9497 19113 1.37 1.0E-07 X4467.1 EST_HUMAN 9661 18222 4.61 1.0E-07 X4467.1 NT 9671 18222 4.61 1.0E-07 X4467.1 NT 9681 18322 4.61 1.0E-07 X4467.1 NT 9852 18730 28025 2.91 9.0E-08 A1734819.1 EST_HUMAN 1054 19508 2.85 2.91 9.0E-08 A11353.1	6103		26132	5.2			EST_HUMAN	1243d08,y1 NCI_CGAP_Brin52 Homo saplens cDNA clone IMAGE:2291339 5'
6424 16286 26447 9.04 1.0E-07 N66081.1 EST_HUMAN 6782 16861 28850 2.81 1.0E-07 P97435 3WISSPROT 6782 16861 28851 2.81 1.0E-07 P97435 3WISSPROT 6782 16861 27865 2.81 1.0E-07 P97435 3WISSPROT 7180 17057 27724 2.37 1.0E-07 BF674624.1 EST_HUMAN 7716 17664 27792 1.28 1.0E-07 AA386311.1 EST_HUMAN 7780 17830 27762 1.56 1.0E-07 AA467.1 NT 9497 18113 25080 3.66 1.0E-07 AA467.1 NT 9497 18113 27765 4.61 1.0E-07 AA467.1 NT 9497 1812 4.61 1.0E-07 AA467.1 NT 9497 18222 4.61 1.0E-07 AA467.1 NT 9497 18222 4.61 1.0E-07 AA467.1 NT 9497 18222 2.86 2.81 0.0E-08 AA467.	6103		26133	5.2			EST_HUMAN	1243d06,y1 NCI_CGAP_Brn52 Homo sapiens cDNA clone IMAGE:2291339 5'
6782 16661 26850 2.81 1.0E-07 P97435 SWISSPROT 6782 16681 28851 2.81 1.0E-07 P97435 SWISSPROT 7180 17057 27246 3.24 1.0E-07 A693576.1 EST_HUMAN 7714 17664 27720 2.37 1.0E-07 A693676.1 EST_HUMAN 7716 17668 27722 1.28 1.0E-07 A4386311.1 EST_HUMAN 7080 17830 25080 3.66 1.0E-07 X4467.1 NT 8497 18113 1.37 1.0E-07 X4467.1 NT 861 18222 4.61 1.0E-07 X4467.1 NT 8520 1832 25680 3.66 1.0E-07 X4467.1 NT 8521 1832 2.66 A7734819.1 EST_HUMAN 8522 18730 28025 3.91 9.0E-08 A173330.2 NT 8922 18730 28025 3.91 9.0E-08 A173330.2 NT 1034 10692 0.88 8.0E-08 B182649.1 EST	6424			9.04	1.0E-07		EST_HUMAN	yw43c07.s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:2454843'
6782 16861 26851 2.81 1.0E-07 P97435 SWISSPROT 7180 17057 27246 3.24 1.0E-07 AA683576.1 EST_HUMAN 7714 17564 27790 2.37 1.0E-07 AA683576.1 EST_HUMAN 7716 17568 27792 1.28 1.0E-07 AA683571.1 EST_HUMAN 7080 17830 25080 3.68 1.0E-07 AA68770.1 EST_HUMAN 8497 19113 1.37 1.0E-07 X64467.1 NT 8520 18392 28656 2.81 8.0E-08 AL6330.2 NT 8521 18730 28025 3.91 9.0E-08 AL6330.2 NT 8922 18730 28025 3.91 9.0E-08 AL6330.2 NT 8922 18730 28025 3.91 9.0E-08 AL6330.2 NT 1034 10692 0.68 8.0E-08 AL6330.2 NT 1034 10692 0.68 8.0E-08 AL6330.2 RT_HUMAN 7074 16951 27144 3.38	6782		26850		1.0E-07	P97435	SWISSPROT	ENTEROPEPTIDASE (ENTEROKINASE)
17657 27246 3.24 1.0E-07 AA693679.1 EST HUMAN 17568 27790 2.37 1.0E-07 BF674524.1 EST HUMAN 17568 27792 1.28 1.0E-07 AA386311.1 EST HUMAN 17650 1.56 1.0E-07 AL163282.2 NT 18533 25060 3.66 1.0E-07 AL163282.2 NT 19113 1.37 1.0E-07 AL163282.2 NT 18322 A.61 1.0E-07 AC4467.1 NT 18322 A.61 1.0E-07 X64467.1 NT 17539 27765 1.67 9.0E-08 AV734819.1 EST HUMAN 18730 28025 3.91 9.0E-08 AV734819.1 EST HUMAN 18730 28026 2.81 9.0E-08 AV734819.1 EST HUMAN 16961 27144 3.8 8.0E-08 BE795469.1 EST HUMAN 16951 27144 3.38 8.0E-08 BE795469.1 EST HUMAN 16951 27144 3.38 8.0E-08 BE795469.1 EST HUMAN 16951 27145 3.38	6782		26851		1.0E-07	P97435	SWISSPROT	ENTEROPEPTIDASE (ENTEROKINASE)
7714 17564 27790 2.37 1.0E-07 BF674624.1 EST HUMAN 7716 17568 27792 1.28 1.0E-07 AA386311.1 EST HUMAN 9364 1853 25060 3.68 1.0E-07 AL163282.2 NT 9467 19113 1.37 1.0E-07 AC467.1 NT 9467 19113 1.37 1.0E-07 X64467.1 NT 9467 18792 27765 1.67 9.0E-08 AL16330.1 EST HUMAN 8520 18392 28656 2.91 9.0E-08 AL16330.1.2 NT 8916 1691 2.65 8.0E-08 AL16330.1.2 NT 8921 1690 3.09 9.0E-08 AL16330.1.2 NT 8922 18730 28025 3.91 9.0E-08 AL16330.1.2 NT 8916 1034 1034 1034 1034 1034 1034 1034 1036 27144 3.38 8.0E-08 AL752367.1 EST HUMAN 7074 16951 27145 3.38 8.0E-08 AL75236	7180		27246		1.0E-07		EST_HUMAN	251e10.s1 Scares_fetal_liver_spleen_1NFLS_S1 Homp sapiens cDNA clone IMAGE:434346 3'
7716 17568 27782 1.28 1.0E-07 AA386311.1 EST HUMAN 7880 17830 1.56 1.0E-07 AL163282.2 NT 9384 18533 25050 3.68 1.0E-07 AL163282.2 NT 9497 19113 1.37 1.0E-07 X64467.1 NT 9661 18222 4.61 1.0E-07 X64467.1 NT 8520 18392 28656 2.91 9.0E-08 AV734819.1 EST_HUMAN 8922 18730 28025 3.91 9.0E-08 AL163301.2 NT 9316 19008 3.09 9.0E-08 AL163301.2 NT 9316 10508 3.09 9.0E-08 AL163301.2 NT 1034 10682 3.09 9.0E-08 AL25495.1 EST_HUMAN 1034 10682 0.68 8.0E-08 BF795469.1 EST_HUMAN 7074 16951 27144 3.38 8.0E-08 AI752367.1 EST_HUMAN 7074 16951 27145 3.38 8.0E-08 AI752367.1 EST_HUMAN 75	77.14	L.,		2.37		BF674524.1	EST_HUMAN	602137714F1 NIH_MGC_83 Homo sapiens cDNA clone IMAGE:4274426 5'
17830 1.56 1.0E-07 AL163282.2 NT 18533 25060 3.66 1.0E-07 BE048770.1 EST_HUMAN 19113 1.37 1.0E-07 X64467.1 NT 18222 4.61 1.0E-07 X51755.1 NT 17539 27765 1.67 9.0E-08 AV734819.1 EST_HUMAN 18730 28025 3.91 9.0E-08 AV734819.1 EST_HUMAN 18730 28025 3.91 9.0E-08 AV734819.1 EST_HUMAN 12601 3.91 9.0E-08 AL163301.2 NT 12601 3.91 9.0E-08 AL251973.1 NT 1265 8.0E-08 BE76449.1 EST_HUMAN 13415 1.43 8.0E-08 BE76449.1 EST_HUMAN 16951 27144 3.38 8.0E-08 BE7649.1 EST_HUMAN 17402 27745 3.38 8.0E-08 BF76367.1 EST_HUMAN 17402 2766 8.0E-08 AI752367.1 EST_HUMAN 17402 27745 3.38 8.0E-08 AI752367.1 EST_HUMAN 17402 2	L	1				AA386311.1	EST_HUMAN	EST185054 Brain IV Homo saplens cDNA
19533 25060 3.66 1.0E-07 RE048770.1 EST_HUMAN 19113 1.37 1.0E-07 X64467.1 NT 18222 4.61 1.0E-07 X51755.1 NT 17539 27765 1.67 9.0E-08 AV734819.1 EST_HUMAN 18392 28658 2.91 9.0E-08 AV734819.1 EST_HUMAN 18730 28025 3.91 9.0E-08 AL163301.2 NT 19008 3.09 9.0E-08 AL163301.2 NT 12651 2.65 8.0E-08 AL163301.2 NT 12652 3.91 9.0E-08 AL251973.1 NT 1265 8.0E-08 BE795469.1 EST_HUMAN 16951 27144 3.38 8.0E-08 BE795469.1 EST_HUMAN 17402 27745 3.38 8.0E-08 AI752367.1 EST_HUMAN 17402 27616 2.89 8.0E-08 AI752367.1 EST_HUMAN 17402 27745 3.38 8.0E-08 AI752367.1 EST_HUMAN 17402 27616 2.89 8.0E-08 AI752367.1 EST_HUMAN	7980	l		1.56		AL163282.2	\ L	Homo sapiens chromosome 21 segment HS21C082
19533 25060 3.66 1.0E-07 RE049770.1 EST_HUMAN 19113 1.37 1.0E-07 X64467.1 NT 18222 4.61 1.0E-07 X51755.1 NT 17539 27765 1.67 9.0E-08 AV734819.1 EST_HUMAN 18392 28656 2.91 9.0E-08 AV734819.1 EST_HUMAN 18730 28025 3.91 9.0E-08 AV734819.1 EST_HUMAN 18730 28025 3.91 9.0E-08 AV734819.1 EST_HUMAN 10963 3.91 9.0E-08 AJ251973.1 NT 10967 0.86 8.0E-08 BJ1352.1 EST_HUMAN 16951 27144 3.38 8.0E-08 BF795469.1 EST_HUMAN 16951 27145 3.38 8.0E-08 BF795469.1 EST_HUMAN 17402 2766 8.0E-08 BF795469.1 EST_HUMAN 16951 27145 3.38 8.0E-08 BF795469.1 EST_HUMAN 17402 2766 8.0E-08 BF795469.1 EST_HUMAN 17402 27745 3.38 8.0E-08 BF795469.1 </td <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td>hr53c11.x1 NCI_CGAP_Kid11 Homo sapiens ¿DNA clone IMAGE:3132212 3' similar to TR.095722 095722</td>								hr53c11.x1 NCI_CGAP_Kid11 Homo sapiens ¿DNA clone IMAGE:3132212 3' similar to TR.095722 095722
19113 1.37 1.0E-07 X64467.1 NT 18222 4.61 1.0E-07 X51755.1 NT 17539 27765 1.67 9.0E-08 AV734819.1 EST_HUMAN 18730 28656 2.91 9.0E-08 AV891052.1 EST_HUMAN 18730 28025 3.91 9.0E-08 AV163301.2 NT 19008 3.09 9.0E-08 AV163301.2 NT 10952 0.86 8.0E-08 AV163301.2 NT 10953 0.86 8.0E-08 BF785469.1 EST_HUMAN 13415 1.43 8.0E-08 BF785469.1 EST_HUMAN 16951 27144 3.38 8.0E-08 AV970683.1 EST_HUMAN 17402 27616 2.89 8.0E-08 AV970683.1 EST_HUMAN 18438 2.39 8.0E-08 AV970683.1 EST_HUMAN 18438 2.39 8.0E-08 AV970683.1 EST_HUMAN	9364			3.66		BE048770.1	EST_HUMAN	DJ1163J1.1;
18222 4.61 1.0E-07 X51755.1 NT 17539 27765 1.67 9.0E-08 AV734819.1 EST_HUMAN 18730 28656 2.91 8.0E-08 AR991052.1 EST_HUMAN 18730 28025 3.91 9.0E-08 AL163301.2 NT 19008 3.09 9.0E-08 AL163301.2 NT 10952 0.28 8.0E-08 AL163301.2 NT 10953 0.28 8.0E-08 BE795469.1 EST_HUMAN 13415 1.43 8.0E-08 BE795469.1 EST_HUMAN 16951 27144 3.38 8.0E-08 AI752367.1 EST_HUMAN 17402 27616 2.89 8.0E-08 AI752367.1 EST_HUMAN 17402 27616 2.89 8.0E-08 AI752367.1 EST_HUMAN 17402 27616 2.89 8.0E-08 AI752367.1 EST_HUMAN 18438 2.39 8.0E-08 AI752367.1 EST_HUMAN	9497			1.37	1.0E-07	X64467.1	LN	H.seplens ALAD gene for porphobilinogen synthase
17539 27765 1.67 9.0E-08 AV734819.1 EST_HUMAN 18730 28656 2.81 9.0E-08 AL93301.2 NT 18730 28025 3.91 9.0E-08 AL163301.2 NT 19008 3.09 9.0E-08 AL163301.2 NT 10952 0.86 8.0E-08 AL163301.2 NT 10953 0.86 8.0E-08 AL951973.1 NT 13415 1.43 8.0E-08 BE795469.1 EST_HUMAN 16951 27144 3.38 8.0E-08 BT752367.1 EST_HUMAN 17402 27616 2.89 8.0E-08 AI752367.1 EST_HUMAN 18438 2.39 8.0E-08 AF283417.1 NT	9661	L		4.61	1.0E-07	X51755.1	NT	Human lambda-immunogiobulin constant region complex (germline)
18392 28656 2.91 9.0E-08 AI891052.1 EST_HUMAN 18730 28025 3.91 9.0E-08 AL163301.2 NT 19008 3.09 9.0E-08 AL251973.1 NT 12671 2.65 8.0E-08 AL251973.1 NT 10952 0.86 8.0E-08 BE795469.1 EST_HUMAN 13415 1.43 8.0E-08 BE795469.1 EST_HUMAN 16951 27144 3.38 8.0E-08 AI752367.1 EST_HUMAN 17402 27616 2.89 8.0E-08 AI752367.1 EST_HUMAN 18438 2.39 8.0E-08 AI752367.1 EST_HUMAN	7689	L	•		9.0E-08	AV734819.1	EST_HUMAN	AV734819 cdA Homo sapiens cDNA clone cdABFB06 5'
18392 28656 2.91 8.0E-08 AI891052.1 EST_HUMAN 18730 28025 3.91 9.0E-08 AL163301.2 NT 19008 3.09 9.0E-08 AJ251973.1 NT 12671 2.65 8.0E-08 AJ251973.1 NT 10952 0.88 8.0E-08 BF795469.1 EST_HUMAN 13415 1.43 8.0E-08 BF795469.1 EST_HUMAN 16951 27144 3.38 8.0E-08 AI752367.1 EST_HUMAN 17402 27616 2.89 8.0E-08 AI752367.1 EST_HUMAN 18438 2.39 8.0E-08 AI752367.1 EST_HUMAN 18438 2.39 8.0E-08 AI752367.1 EST_HUMAN								wn30a07.x1 NCI_CGAP_Gas4 Homo sapiens cDNA clone IMAGE:2446932 3' similæ to contains OFR.t2
18730 28025 3.91 9.0E-08 AL163301.2 NT 19008 3.09 9.0E-08 AJ251973.1 NT 12671 2.65 8.0E-08 AJ251973.1 NT 10952 0.88 8.0E-08 BE795469.1 EST_HUMAN 13415 1.43 8.0E-08 BE795469.1 EST_HUMAN 16951 27144 3.38 8.0E-08 AI752367.1 EST_HUMAN 17402 27616 2.89 8.0E-08 AI752367.1 EST_HUMAN 18438 2.39 8.0E-08 AI752367.1 EST_HUMAN	8520				9.0E-08	AI891052.1	EST_HUMAN	OFR repetitive element;
1900B 3.09 9.0E-08 AJ251973.1 NT 12671 2.65 8.0E-08 AJ251973.1 EST_HUMAN 10952 0.88 8.0E-08 BE795469.1 EST_HUMAN 13415 1.43 8.0E-08 BE795469.1 EST_HUMAN 16951 277144 3.38 8.0E-08 AI752367.1 EST_HUMAN 17402 27616 2.89 8.0E-08 AI752367.1 EST_HUMAN 17402 27616 2.89 8.0E-08 AI752367.1 EST_HUMAN 18438 2.39 8.0E-08 AI752347.1 NT	8922			3.91	9.0E-08	AL 163301.2	NT	Homo saplens chromosome 21 segment HS21C101
12671 2.65 8.0E-08 Al911352.1 EST_HUMAN 10952 0.88 8.0E-08 BE795469.1 EST_HUMAN 13415 1.43 8.0E-08 BE795469.1 EST_HUMAN 16951 27144 3.38 8.0E-08 Al752367.1 EST_HUMAN 17402 27616 2.89 8.0E-08 Al752367.1 EST_HUMAN 18438 2.39 8.0E-08 Al752367.1 EST_HUMAN 18438 2.39 8.0E-08 Al752347.1 INT_HUMAN	9316			3.09	9.0E-08	AJ251973.1	NT	Homo sapiens partial steerin-1 gene
10952 0.88 8.0E-08 BE795469.1 EST_HUMAN 13415 1.43 8.0E-08 BE795469.1 EST_HUMAN 16951 27144 3.38 8.0E-08 AI752367.1 EST_HUMAN 16951 27145 3.38 8.0E-08 AI752367.1 EST_HUMAN 17402 27616 2.89 8.0E-08 AW970693.1 EST_HUMAN 18438 2.39 8.0E-08 AF253417.1 NT	294			2.65	8.0E-08	AI911352.1	EST_HUMAN	wd16b06.x1 Soares_NFL_T_GBC_S1 Hamo supiens cDNA clane IMAGE:2328273 3'
13415 1,43 8.0E-08 BE795469.1 EST_HUMAN 16961 27144 3.38 8.0E-08 AI752367.1 EST_HUMAN 16951 27445 3.38 8.0E-08 AI752367.1 EST_HUMAN 17402 27616 2.89 8.0E-08 AW970693.1 EST_HUMAN 18438 2.39 8.0E-08 AF253417.1 NT	1034			0.88	8.0E-08	BE795469.1	EST_HUMAN	801590133F1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:3943976 5'
16961 27144 3.38 8.0E-08 AI752367.1 EST_HUMAN 16951 27145 3.38 8.0E-08 AI752367.1 EST_HUMAN 17402 27616 2.89 8.0E-08 AW970693.1 EST_HUMAN 18438 2.39 8.0E-08 AF253417.1 NT	3498	Ш		1,43	8.0E-08		EST_HUMAN	601590133F1 NIH_MGC_7 Homo saplens cDNA clone IMAGE:3943976 5'
16951 27145 3.38 8.0E-08 AI752367.1 EST_HUMAN 17402 27616 2.89 8.0E-08 AW970693.1 EST_HUMAN 18438 2.39 8.0E-08 AF283417.1 NT	7074		27144	88		A[752367.1	EST HUMAN	cn15c02.x1 Normal Human Trabecular Bone Cells Homo sapiens cDNA clone NHTBC cn15c02 random
16851 2745 3.38 8.0E-08 AI752367.1 EST_HUMAN 17402 27616 2.89 8.0E-08 AW970693.1 EST_HUMAN 18438 2.39 8.0E-08 AF253417.1 NT								
17402 27616 2.89 8.0E-08 AW970693.1 EST_HUMAN 18438 2.39 8.0E-08 AF263417.1 NT	7074						EST_HUMAN	cn15c02.x1 Normal Human Trabecular Bone Cells Homo sapiens cDNA clone NHTBC_cn15c02 random
18438 2.39 8.0E-08 AF253417.1 NT	7551			2.89	8.0E-08	AW970693.1	EST_HUMAN	EST382776 MAGE resequences, MAGK Homo sepiens cDNA
	8570			2.39	8.0E-08	AF253417.1	LZ.	Homo saplens microsomal epoxide hydrolase (EPHX1) gene, complete cds

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Probe SEQ ID NO:	SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
73	10057	19874	3.27	7.0E-08	8 Q02357	SWISSPROT	ANKYRIN 1 (ERYTHROCYTE ANKYRIN)
1338	3 11244	21102	12.71	7.0E-08 X04809.		IN	Rat mRNA for ribosomal protein L31
3528	3 13444	23240	1.09	7.0E-08 P15305		SWISSPROT	DYNEIN HEAVY CHAIN (DYHC)
3528	13444		1.09	7.0E-08 P15305		SWISSPROT	DYNEIN HEAVY CHAIN (DYHC)
8189	18075		4.63	7.0E-08	7.0E-08 A1535743.1	EST_HUMAN	cong3.P11.A5 conorm Homo sapiens cDNA 3
8926	18734		9.6			TN	Rattus norvegicus Munc13-1 mRNA, complete cds
9770	13444		4.54	7.0E-08 P15305		SWISSPROT	DYNEIN HEAVY CHAIN (DYHC)
9770	13444	23241	4.54	7.0E-08 P15305		SWISSPROT	DYNEIN HEAVY CHAIN (DYHC)
9834	19334		1.89	7.0E-0	8 AJ131016.1	NT	Homo saplens SCL gene locus
88	10729	20569	2.84	6.0E-08	6.0E-08 AL163248.2	NT	Homo sapiens chromosome 21 segment HS21C048
800	10729	20570	2.84	8.0E-08	6.0E-08 AL163248.2	IN	Homo saplens chramosame 21 segment HS21C048
2312	12193		1.73	6.0E-08	6.0E-08 BE144398.1	EST_HUMAN	MR0-HT0166-191199-004-509 HT0166 Homo sapiens cDNA
4165	14055		66.0	80-30.9	6.0E-08 AL163248.2	۲	Homo saplens chromosome 21 segment HS21C048
							RETROVIRUS-RELATED POL POLYPROTE:IN CONTAINS: REVERSE TRANSCRIPTASE;
8717		28818	2.28			SWISSPROT	ENDONUCLEASE]
8819	18632		1.74	6.0E-08		IN	Homo sapiens chromosome 21 segment HS21 C009
79	10063	19880	3.08		5.0E-08/AL163303.2	LN	Homo sapiens chromosome 21 segment HS21C103
							nh03b09.s1 NCI_CGAP_Thy1 Homo sapiens cDNA clone IMAGE:943193 similar to contains Alu repetitive
2189			1.97	5.0E-08	51.1	EST HUMAN	element,
5178	15042	L	1.12	5.0E-08 Q06278		SWISSPROT	ALDEHYDE OXIDASE
8057	18839		4.38	5.0E-08 P06681		SWISSPROT	COMPLEMENT C2 PRECURSOR (C3/C5 CONVERTASE)
9245	5 18958	25317	1.94	5.0E-08	5.0E-08 AW851878.1	EST_HUMAN	QV0-CT0225-131099-034-a12 CT0225 Homo saplens cDNA
1724	11625	21493	1.1	4.0E-08 P25723		SWISSPROT	DORSAL-VENTRAL PATTERNING TOLLOID PROTEIN PRECURSOR
1724	11625		1.1	4.0E-08		SWISSPROT	DORSAL-VENTRAL PATTERNING TOLLOID PROTEIN PRECURSOR
7278	3 17155		1.41	4.0E-08	4.0E-08 L42571.1	TN	Cricetulus griseus ribosomal transcription factor (UBF2) mRNA, complete cds
							an22d10.x1 Gessler Wilms tumor Homo sapiens cDNA done IMAGE:1699411 3' similar to contains Alu
7838	17788	28030	4.17	4.0E-08	4.0E-08 A1050027.1	EST_HUMAN	repetitive element;contains element MER22 repetitive element;
8274	ı		1.79	4.0E-08		IN	Homo sapiens mRNA for UGA suppressor tRNA-associated antigenic protein (tRNA48 gene)
8439	18313		3.66	4.0E-08	4.0E-08 BF692493.1	EST_HUMAN	602248024F1 NIH_MGC_62 Homo sapiens cDNA clone IMAGE:4333300 5'
8439	18313	28571	3.68	4.0E-08	4.0E-08 BF692493.1	EST_HUMAN	602248024F1 NIH_MGC_62 Homo sapiens cDNA clone IMAGE:4333300 5'
	<u> </u>						zd65g03.r1 Soares_fetal_heart_NbHH19W. Homo sepiens cDNA clone IMAGE:345566 5' similar to contains
906	19880		1.54	4.0E-08	4.0E-08 W76159.1	EST HUMAN	L1.t1 L1 repetitive element;
9703	19252		1.46		4.0E-08 AI343353.1	EST HUMAN	tb95a11.x1 NCI_CGAP_Co16 Homo sapiens cDNA clone IMAGE:2062076 3' similar to conteins MER18.b3 MER18 MER18 repetitive element ;
	ı			l			

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zw48f07.r1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA clone IMAGE:773317 5' similar to contains th99h09.x1 Soares_NSF_F8_9W_OT_PA_P_S1 Homo saplens cDNA clone IMAGE:2126273 3' similar to TR:Q13537 Q13537 MER37 TRANSPOSABLE ELEMENT, COMPLETE CONSENSUS SEQUENCE.; bb79e10.y1 NIH_MGC_10 Homo sapiens oCiNA clone IMAGE:3048670 5' similar to TR:092158 Q92158 SYNTAXIN 17.; nw64h01.s1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:1251409 3' similar to contains L1.t3 L1 Y902f04.r1 Soares Infant brain 1NIB Homo sapiens cDNA clone IMACE:30948 5' similar to contains Atu Homo seplens shox gene, alternatively spliced products, complete cds aa28c07.r1 NCI_CGAP_GCB1 Homo sepiens cDNA clone IMAGE:814380 5' similar to contains L1.t2 L1 he17h08.x2 NCf_CGAP_CML1 Homo sapiens cDNA clane IMAGE:2919327 3' similar to contains Alu zq45d06.r1 Stratagene hNT neuron (#937233) Homo sapiens cDNA clone IMAGE:632849 5 2445d05.r1 Stratagene hNT neuron (#637233) Homo sapiens cDNA clone IMAGE:632649 ab02g06.s1 Stratagene fetal retina 937202 Homo saplens cDNA clone IMAGE:839674 3* nas32e09.x1 NCI_CGAP_Kid11 Homo saplens cDNA clone IMAGE:3257869 3* qs76f11.y5 NCI_CGAP_Pr28 Homo sapiens cDNA clone IMAGE:1944045 6' Homo sapiens chromosome 21 segment HS21C046 x87108.x1 NCI_CGAP_Lu26 Homo seplens cDNA clone IMAGE:2767139 3 MR0-OT0080-240200-001-g08 OT0080 Homo sepiens cDNA 801155321F1 NIH_MGC_21 Homo sepiens cDNA clone IMAGE:3138893 5 601570463F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3845199 5 49411 X1 NCI_CGAP_HN11 Homo sapiens cDNA clone IMAGE:2743149 Alu repetitive element; contains element MER15 repetitive element Top Hit Descriptor MR0-OT0080-240200-001-g08 OT0080 Homo saplens cDNA WNT-14 PROTEIN PRECURSOR RC3-ST0197-161099-012-b03 ST0197 Hamo sapiens cDNA Gallus gallus Dach2 protein (Dach2) mRNA, complete cds Homo saplens chromosome 21 segment HS210047 WNT-14 PROTEIN PRECURSOR Single Exon Probes Expressed in Heart Sheep His-tRNA-GUG repetitive element; repetitive element repetitive element EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN Top Hit Database EST HUMAN EST_HUMAN EST_HUMAN EST_HUMAN NT EST_HUMAN EST_HUMAN Source EST HUMAN EST HUMAN HUMAN EST HUMAN EST_HUMAN EST HUMAN SWISSPRO EST K 눋 5 Fop Hit Acession 3.0E-08 BE018348.1 3.0E-08 AI792737.1 3.0E-08 AL163246.2 2.0E-08 | AF198349.1 2.0E-08 | AW886438.1 2.0E-08 | AW886438.1 2.0E-08 | BE280477.1 2.0E-08 | AL163247.2 2.0E-08 AW 302996.1 3.0E-08 AA191195.1 3.0E-08 AA191195.1 3.0E-08 AI436352.1 2.0E-08 AA425598.1 AW270271.1 ģ 2.0E-08 AW572881.1 2.0E-08 BE734871.1 2.0E-08 AA731948.1 2.0E-08 AA490121.1 2.0E-08 BF589904.1 2.0E-08 AA459040.1 3.0E-08 R18420.1 2.0E-08 K00216.1 2.0E-08 O42280 K00216.1 **U82668.1** 2.0E-08 O42280 2.0E-08 AW8136 2.0E-08 U82688.1 (Top) Hit BLAST E Most Simila 20E-08 Value 7.24 99 11.6 3.58 2.97 10.93 10.93 37.42 Expression 7.97 1.48 1.83 1.66 3.89 1.42 2.16 6.02 6.02 3.35 288 11.62 ب 24812 ORF SEQ 24884 26478 20396 20397 ÖNO 21082 22140 22902 22901 23669 28082 26791 15048 SEQ ID 15120 15367 16312 16416 18819 10194 10431 11228 1917 11608 12248 10581 10898 ġ 13096 12371 13098 13893 14203 14756 16602 18789 SEO ID 5184 5184 8446 6163 6558 6451 189 1319 1707 9027 233 488 4 4 974 2368 2496 3171 3783 6722 ö 3171 3986 4876 8895 4305

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	_	_				_	_	_	_			_	_	_	_		_	ψ,		_		_		_	_	_	_			_
Top Hit Descriptor	POL POLYPROTEIN (CONTAINS: REVERSIE TRANSCRIPTASE ; ENDONUCLEASE)	PM2-HT0130-150999-001-f12 HT0130 Homo saplens cDNA	TCBAP105232 Pediatric pre-B cell acute lymphoblastic leukemia Baylor-HGSC project=TCBA Homo saniens cDNA clane TCBAP5332	TCBAP1D5232 Pediatric pre-B cell acute (ymphoblastic leukemia Baylor-HGSC project=TCBA Homo	sapiens cDNA done TCBAP5232	Homo sapiens chromosome 21 segment HS21C080	Homo sapiens hyperion gene, exons 1-50	ot35a05.s1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1618736 3'	Homo sapiens major histocompatibility locus class III region	Human lambda-immunoglobulin constant region complex (germline)	Human lambda-immunoglobulin constant region complex (germline)	MRA-ST0240-240700-013-g04 ST0240 Homo sapiens cDNA	Homo sapiens chromosome 21 segment HS21C079	Homo sapiens chromosome 21 segment HS21,0079	RC5-BN1058-270400-031-C06 BN1058 Homo sapiens cDNA	qd42e07.x1 Soares_fetal_heart_NbHH19W Homo sapiens cDNA clone IMAGE:1732164.3' similar to contains MSR1.t1 MSR1 repetitive element;	OMO-NN1004-100300-273-e06 NN1004 Home sapiens cDNA	op74d08.s1 Soares_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:15825753'	Homo sapiens DNA for 3-ketoacyk-CoA thiolase beta-subunit of mitochondrial trifunctional protein, exon 2, 3	Homo sapiens gene for enteric smooth muscle gamma-actin, exon 2, 3	Human lysosomal membrane glycoprotein-2 (L∤MP2) gene, 5' end and flanking region	601111173F1 NIH_MGC_16 Homo saplens cDNA clone IMAGE:3351834 5'	DKFZp434C0514_r1 434 (synonym: htes3) Homo saplens cDNA clane DKFZp434C0514 5'	n17a11.s1 NCI_CGAP_HSC1 Homo sapiens cDNA clone IMAGE:1040924 similar to contains L1.t2 L1	iejoeunve etement	PM1-H10527-160200-001-h05 H10527 Homo sapiens cDNA	xn85h08.x1 Soares_NFL_T_GBC_S1 Homo expiens cDNA clone IMAGE:2701311 3'	Homo sapiens fibroblast growth factor receptor 3 (achondroplasia, thanatophoric dwarfism) (FGFR3) mRNA	Homo sapiens testis-specific kinase substrate (TSKS) gene, complete cds	RC2-HT0252-120200-014-h10 HT0252 Homo sapiens cDNA
Top Hit Database Source	SWISSPROT	EST HUMAN	NAMI H TAR	N	EST_HUMAN	LN	Z	EST_HUMAN	LZ L	LN	LN	EST_HUMAN	LN.	Z	EST_HUMAN	EST HUMAN	EST HUMAN		TN	LN	INT	EST_HUMAN	EST_HUMAN	100	ES HOMAIN	EST_HUMAN	EST HUMAN	TN	NT	EST_HUMAN
Top Hit Acessian No.	P31792	BE141959.1	RE 246844 1		BE246844.1		AJ010770.1	AID15304.1	AF044083.1	X51755.1	X51755.1	BF375398.1	AL163279.2	AL163279.2	BE012076.1	Al183500.1	AW900159.1	AA938892.1	D86842.1	D00649.1	1.09709.1	BE254850.1	AL040439.1		AA55/940.1	BE169421.1	AW195784.1	4503710 NT	AF200923.2	BE149264.1
Most Similar (Top) Hit BLAST E Value	1.0E-08	1.0E-08	4 OF - DR	20-10-1	1.0E-08	1.0E-08	1.0E-08	1.0E-08	1.0E-08	1.0E-08	1.0E-08	1.0E-08	60-30.6	9.0E-09	8.0E-09	8.0E-09	8.0E-09	8.0E-09	7.0E-09	7.0E-09	7.0E-09	7.0E-09	6.0E-09	00 100	9.0E-08	6.0E-09	8.0E-09	6.0E-09	6.0E-09	5.0E-09
Expression Signal	1.13	2:92	1 24	rai.	1.24	1.43	4.05	1.84	3.66	2.01	6.3	1.76	2.98	2.98	1.07	6.12	2.71	3.06	1.68	0.99	3.27	1.84	1.06		1.00	4.44	8.19	2.26	4.06	3.09
ORF SEQ ID NO:	21257		7.88.1		22882	24818		27045	28773				23823	ļ		26328				24236	27403					24562		27370		21157
Exon SEQ ID NO:		11897	13080	1	13080	15054	15358	16852	18498	19074	19316	19518	14049	14049	13381	16171	16546	l	13469	14450	17203	17700	11996		ļ	1	15228	17170	17748	11299
Probe SEQ ID NO:	1483	2002	24 55		3155	5191	5438	6975	8633	9434	9804	9894	4149	4149	3444	6307	9999	7186	3555	4558	7335	7850	2107		1486	4806	6305	7294	7896	1394

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Table 4
Single Exon Probes Expressed in Heart

Probe Exon ORF SEQ ID ID NO: NO: NO: NO: NO: NO: NO: NO: NO: NO:	֡				
15764 17661 10452 10873 11358 12259 1226 1426 1426 14305 14305 14305 14305 14305 14305 14305 14305 14305 14305 14305 14305 14305 16724 11144 11144 11144 11144 11144	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
17661 10452 11358 1259 12759 12780 1428 14228 14228 14305 14305 14305 14305 14305 14305 14305 14305 14305 14305 17734 11144 11144 11144 11144 11144 11144 11144	1.92	5.0E-09 A	AA359454.1	EST_HUMAN	EST68746 Fetal lung II Homo sapiens cDNA 5' end
10873 11358 12259 12259 12280 13208 14228 14305 14305 14305 14305 11144 11144 11144 11144 11144	1 2.69	5.0E-09	5.0E-09 AW799667.1	EST_HUMAN	PM2-UM0053-240300-005-c09 UM0053 Homo sapiens cDNA
12259 12259 12259 12280 13208 14228 1428 14	1.68	4.0E-09	AL163282.2	NT	Homo sapiens chromosome 21 segment HS2 i C082
12259 12259 12259 12380 12482 14228 14305	2.31	4.0E-09	AL163285.2	LN	Homo sapiens chromosome 21 segment HS21C085
12259 12184 12380 12482 14228 14305 14305 14305 14305 14305 14305 14305 14305 14305 14305 14305 14305 14305 14305 14305 14305 14305 16261	2 0.95	4.0E-09	9558718 NT	LN	Homo sapiens hypothetical protein (AF038169), mRNA
12184 12380 12482 13208 14228 14305 14305 14305 14305 14305 14305 14305 14305 14305 14305 14305 16261	1 6.36	4.0E-09	4.0E-09 AA350878.1	EST_HUMAN	EST58385 Infant brain Homo sapiens cDNA 5' end similar to similar to heat shock protein, 90 kDa
12380 12482 13208 14226 14305 17734 18261 10724 11144 11144 11144 111641			7 000000	10 TO TO TO	hu09e09.x1 NCI_CGAP_Lu24 Homo sepiens 5DNA clone IMAGE:3168120 3' similar to contains MER18.t3
12380 13208 14208 14208 14305 17734 18261 10724 11144 11541	3.82	3.05-09 12	BE222239.1	ESI_HUMAN	MENTO repetitive element
12492 13208 14228 14305 14734 18281 18281 10724 11144 11541 12162	1.26	3.05-09	BE222239.1	EST HUMAN	hu09e09.x1 NCI_CGAP_Lu24 Homo sapiens ≎DNA clone IMAGE:3166120 3' similar to contains MER18.t3 MER18 repetitive element ;
13208 14226 14305 17734 18261 10724 11144 11541 12162			P23249	SWISSPROT	PROTEIN MOV-10
13208 14226 14305 17734 18261 10724 11144 11144 11541					hu09e09.x1 NCI_CGAP_Lu24 Homo sapiens ¿DNA clone IMAGE:3168120 3' similar to contains MER18.t3
14226 14305 17734 16261 10724 11144 11541 12162	1.1	3.0E-09 B		EST_HUMAN	MER18 repetitive element;
14305 17734 18281 18281 10724 11144 11541 12162	3.22	3.0E-09 AF175325	.1	NT	Homo sapiens eukaryotic initiation factor 4AI (EIF4A1) gene, partial ods
17734 18261 10724 11144 11541 12162		3.0E-09		SWISSPROT	258.1 KDA PROTEIN C210RF5 (KIAA0933)
18261 10724 11144 11541 12162	1.73	3.0E-09 A		FA	Homo sapiens chromosome 21 segment HS21C047
18261 10724 11144 11541 12162 13769	3.8	3.0E-09	F109943.1	EST_HUMAN	7172c08.x1 Soares_NSF_F8_9W_OT_PA_P_S1 Homo saplens cDNA clone IMAGE:3527030 3'
10724 11144 11541 12162 13769	1 3.8	3.0E-09	3.0E-09 BF109943.1	EST_HUMAN	7172c08.x1 Sogres_NSF_F8_9W_OT_PA_P_S1 Homo sapiens cDNA clone IMAGE:3527030 3'
11144 11541 12162 13769	83.0	2.0E-09		NT	H.sapiens PADPRP-I gene for NAD(+) ADP-ribosytransferase
11541 12162 13769	5 5.23	2.0E-09 A	AL163284.2	TN	Homo saplens chromosome 21 segment HS21C084
12162 13769	10.52	2.0E-0	9 AL118573.1	EST_HUMAN	DKFZp781B1710_r1 761 (synonym: hamy2) Homo sapiens cDNA clone DKFZp761B1710 5'
13769		2.0E-0(QOY3R5	SWISSPROT	258.1 KDA PROTEIN C210RF5 (KIAA0933)
		2.0E-09	060241	SWISSPROT	BRAIN-SPECIFIC ANGIOGENESIS INHIBITOR 2 PRECURSOR
_					zx63h06,r1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA clone IMAGE:796187 5' similar to contains
16266	ı	2.0E-09		EST_HUMAN	Alu repetitive element;
	1.37	2.0E-09	2.0E-09 AJ271735.1	NT	Homo sapiens Xq pseudoautosomal region; segment 1/2
	3 2.11	2.0E-09		L	Homo sapiens chromosome 21 segment HS21:0048
9578 10724	11.53	2.0E-09		NT	H.saplens PADPRP-I gene for NAD(+) ADP-ribosyltransferase
					nc11c02.r1 NCI_CGAP_Pr1 Homo saplens cDNA clone IMAGE:1007810 similar to contains Alu repetitive
19749	1.62	2.0E-09 A	2.0E-09 AA226070.1	EST_HUMAN	element;
		1.0E-09	5031624 NT	NT	Homo sapiens CCAAT-box-binding transcription factor (CBF2) mRNA
11009		1.0E-08	5031624 NT	NT	Homo sapiens CCAAT-box-binding transcription factor (CBF2) mRNA
1616 11520	0.95	1.0E-08	AJ229041.1	NT	Homo sapiens 959 kb contig between AML1 and CBR1 on chromosome 21q22, segment 1/3

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
2453	12330		0.94	1.0E-09	1.0E-09 Al356086.1	EST_HUMAN	qy64e11.x1 NCI_CGAP_Bm25 Homo saplens cDNA clone IMAGE:2016812.3' similar to contains MER12.t2 MER12 repetitive element;
2860	12788	22580	1.51	1.0E-09	1.0E-09 U80017.1	١	Homo sapiens basic transcription factor 2 p44'(bt/2p44) gene, partial cds, neuronal apoptosis Inhibitory protein (naip) and survival motor neuron protein (smn) genes, complete cds
2895	12822	22814		1.0E-09	1.0E-09 M28699.1	NT	Homo sapiens nucleolar phosphoprotein B23 (NPM1) mRNA, complete cds
2895	12822	22815		1.0E-09	1.0E-09 M28699.1	NT	Hano sapiens nucleolar phosphoprotein B23 (NPM1) mRNA, complete cds
2949	12876		1.23	1.0E-09 P11799		SWISSPROT	MYOSIN LIGHT CHAIN KINASE, SMOOTH MUSCLE AND NON-MUSCLE ISOZYMES (MLCK) [CONTAINS: TELOKIN]
3002	[27.22	0.78	1.0E-09	40.1	Г	801058602F1 NIH_MGC_10 Homo sapiens cDNA clone IMAGE:3445177 5'
4692	l			1.0E-09			zh35b03.s1 Soares, pineal, gland, N3HPG Homo saplens cDNA clone IMAGE:414029 3' similar to contains. Alu repetitive element;contains element MER22 repetitive element;
5560	15476	<u> </u>	1.37	1.0E-09	1.0E-09 U07000.1	LN	Human breakpoint cluster region (BCR) gene, complete cds
5736	15844	25749	3.21	1.0E-09 P26694		SWISSPROT	CIRCUMSPOROZOITE PROTEIN PRECURSOR (CS)
7911	17761		3.1	1.0E-09	33.2	LN	Homo saplens chromosome 21 segment HS210083
9478	19689		2.14	1.0E-09	11418127 NT		Homo saplens GTP binding protein 1 (GTPBP1), mRNA
1287	11195	21048	1.8	9.0E-10	9.0E-10 AW867740.1	EST_HUMAN	MR0-SN0040-050500-002-c07 SN0040 Home sapiens cDNA
2803	12733	22532	6.03	9.0E-10	0 A1870071.1	EST_HUMAN	we78h03.x1 Soares_Dieckgraefe_colon_NHCb Homo sapiens cDNA clone IMAGE:2347253 3' similar to SW:RL29_HUMAN P47914 60S RIBOSOMAL PROTEIN L29 ;contains element PTR5 repetitive element;
6087	16032	26172	4.63	9.0E-10	0 AI452982.1		ij48b09.x1 Soares, NSF F8_9W_OT_PA_P_\$1 Home sapiens cDNA clone IMAGE:2144537 3' similar to TR:000372 000372 PUTATIVE P150.;
141	1		9.43	8.0E-10		TN	Homo sapiens MCM4 (MCM4) and DNA-PKcs (PRKDC) genes, partial cds
3300	13222	23022	78.0	8.0E-10	0 BE080748.1	EST_HUMAN	QV1-BT0631-150200-071-f01 BT0631 Homo sapiens cDNA
4106	14006	23782	2.82	8.0E-10	0 AA376832.1	T_HUMAN	EST89664 Small intestine I Homo capiens cDNA 6' end
7725	17575		2.32	8.0E-10	0 U36308.2	ĽΝ	Homo sapiens lens major intrinsic protein (MIP) gene, complete cds
3008	18808	29100	2.31	8.0E-10	0 AL163280.2	NT	Homo sapiens chromosome 21 segment HS210080
989	10619		12.45	7.0E-10	7706225 NT	ZT.	Homo sapiens TPA inducible protein (LOC51686), mRNA
989	10619	20445	12.45	7.0E-10	7708225 NT	Ę	Homo saplens TPA inducible protein (LOC51836), mRNA
1605	11510		1.87	7.0E-10	0 Q13342	SWISSPROT	LYSP100 PROTEIN (LYMPHOID-RESTRICTED HOMOLOG OF SP100)
1974	11867		1.01	7.0E-10	0 P08548	SWISSPROT	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG
2512	12388		16.88	7.0E-10 P08547		SWISSPROT	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG
3049		22768	2.65	7.0E-10	0 X00856.1	NT	H.sapiens DHFR gene, exon 3
5754	П	Ш	3.98	7.0E-10	20.1	EST_HUMAN	EST51247 Gall bladder II Homo sapiens cDNA 5' end
6514	16373		1.39	7.0E-10	0 P35084	SWISSPROT	DNA-DIRECTED RNA POLYMERASE II LARGEST SUBUNIT
	į						

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Expression Signal (Top) Hit Acession Signal Top Hit Acession Signal Top Hit Descriptor Source Top Hit Descriptor	Homo sapiens ASCL3 gene, CEGP1 gene, C11orf14 gene, C11orf16 gene and C11orf17 gene and C11orf17 gene and C11orf17 gene	1.47 6.0E-10 A1424405.1 [EST_HUMAN t002d07.x1 NCI_CGAP_Pt28 Homo sapiens cDNA clone IMAGE::2095021 3'	AW853719.1 EST_HUMAN	1.79 6.0E-10 AW971823.1 EST HUMAN EST384012 MAGE resequences, MAGL Homo sepiens cDNA	BE699410.1 EST_HUMAN	4.5 5.0E-10 AL046804.1 EST_HUMAN DKFZp434N219_r1 434 (synonym: htes3) Homo sepiens cDNA clone DKFZp434N219 5'	1.48 5.0E-10 Q01033 SWISSPROT HYPOTHETICAL GENE 48 PROTEIN	BF105159.1 EST_HUMAN	P34678 SWISSPROT	1.79 5.0E-10 P34678 SWISSPROT HYPOTHETICAL 67.9 KD PROTEIN ZK688.8 IN CHROMOSOME III	qg09f09.x1 Soares_placenta_8t09weeks_2NbHP8to9W Homo saplens cDNA clone IMAGE:1759049 3: 0.99 4.0E-10 A)221083.1 EST HUMAN similar to contains LTR8 to LTR8 repetitive element;		AW 594709.1 EST_HUMAN	6.09 4.0E-10 AL163303.2 INT Homo sepiens chromosome 21 segment HS21C103	Homo eapiens mannosidase, bota A, lysosome! (MANBA) gene, and ubiquitin-conjugating enzyme E2D 3 (UBE2D3) genes, complete cds		7.7 4.0E-10 AF003528.1 NT regions	1.72 3.0E-10 N3813.1 EST HUMAN L1.11 L1 repetitive element:	3.0E-10 AY005150.1 NT	1.1 3.0E-10 AL163203.2 NT Homo sapiens chromosome 21 segment HS21C003	1.1 3.0E-10 AL163203.2 NT Homo sepiens chromosome 21 segment HS21C003	3.0E-10 P20350 SWISSPROT	3.0E-10 BE302970.1 EST_HUMAN	1.31 3.0E-10 AV743302.1 EST_HUMAN AV743302 CB Hamo sapiens cDNA clone CBł BGD08 5'			H87208.1 EST_HUMAN	AW850731.1 EST_HUMAN	3.0E-10 AW850731.1 EST_HUMAN	2 AC 40 TERSON 4 TECT III MAN In 14012 11 Statement line (#037210) Home caniene CDNA Clerk MARE 80308 5'
Most Similar (Top) Hit BLAST E Value	6.0E-10	6.0E-10	6.0E-10	6.0E-10	6.0E-10	5,0E-10	5.0E-10	5.0E-10	5.0E-10	5.0E-10	4.0E-10		4.0E-10	4.0E-10	4.0E-10		4.0E-10	3.0E-10	3.0E-10	3.0E-10	3.0E-10	3.0E-10	3.0E-10	3.0E-10	3.0E-10		3.0E-10	3.0E-10	3.0E-10	
Exan . ORF SEQ ESQ ID NO:	10821 20666	12511 22402	14522	18865	19597	10675	13347 23152	16197	17368 27570	17366 27571	10087		11846 21732	12401 22292	16125 26278		18310	10822 20668		14330 24117	14330 24118	15672 25779	15739 25851	16420 26599	16420 26600				17110 27303	
Probe SEQ ID SI NO:	895	2644	4634		9947	747	3430		1	7496	106	L	1951	2527	6259	1	8436	897	1_	4435	4435	Ĺ			6562	[- 1	1	1	

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Table 4:
Single Exon Probes Expressed it

Probe	Ä			Most Similar		igle Exon Pro	Single Exon Probes Expressed in Heart
SEQ ID	0)	ORF SEQ ID NO:	Expression Signal	(Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
8026	LI	Ll	1.54	3.0E-10	AA769294.1	EST HUMAN	nzibani si Nci Coab ocea
9737		25227	2.03	3.0E-10	BE179517 1	FOT HIMAN	12 UTAGE 4147504 105
32	10019	19814	1.43	2.0E-10	P48988	SWISSPBOT	MA IOD CENTRONED A LEGIS TO A 10018 HOMO Saplens cDNA
32	10019	19815	1.43	2.0E-10	P48988	SWISSPROT	MA 10R CENTROMEDE AUTOMITIGEN B (CENTROMERE PROTEIN B) (CENP-B)
1855	11751		5.91	2.0E-10		F	Homo septimical and index (CENTROMERE PROTEIN B) (CENP-B) None septimical and actor 2 p44 (bf2p44) gene, partial cds, neuronal apoptosis inhibitory
6647	15463		2.41	2.0E-10		SWISSPROT	Protein (naty) and survival motor heuron protein (smn) genes, complete cds
5787	15693	25801	1.74	2000	A EDOMANT A	į	Homo saplens cytochrome P450 polypeptide 43 (CYP3A43) gene, partial cds; cytochrome P450 polypeptide 4 (CYP3A4) and cytochrome P450 polypeptide 7 (CYP3A7) genes, complete cds; and cytochrome P450 polypeptide
6367	Ĺ	26389	5.79		BE791082 1	EST HIMAN	Polypeptide 5 (CYP3A5) gene, partial cds
1492	11397	 -	3.58		AW867767 4	ECT LANKE	VIOLOGOZOGI I MIH. MGC. / Homo saplens cDNA clone IMAGE:3940824 5
1589	11493	21353	3.14		AV652123 1	EST LIMAN	MKN-SN0U36-2603U0-001-f01 SN0038 Homo sapiens cDNA
2537	12411	-	2.4	1.0E-10	AW852001 1	EST LI MANN	Average Action Sapiens cDNA clone GLCCXA113'
3456	13372	23178	0.89	1.0E-10	AW832912 1	EST HIMAN	0.00 - 0.
3770	13411		0.91		T	EST LIMAN	TALEST CONTROL OF THE
			 -			NICH LICIANA	UNITEPASANTST/_T1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434N1317 5
3333	13842	+	5.44	1.0E-10	AF213884.1	N	nomo sapiens nuclear factor of kappa light polypeptide gene enhancer in B-cells 1 (NFKB1) gene, complete cds
4036	13939	23716	4.61	1.0E-10	U52111.2	F	Homo saplens X28 region near ALD locus containing dual specificity phosphatase 9 (DUSP9), ribosomal protein L18a (RPL18a), Ca2+/Calmodulin-dependent protein kinase I (CAMKI), creatine transporter (CRTR), CDM protein (CDM), advanced into the contained of the con
							Homo saplens X28 region near ALD locus containing dual specificity phosphatasa 9 (DJ ISP9), ribreomal
4038	13939	23717	4.51	1.0E-10	U52111.2	Ę	protein L18a (RPL18a), Ca2+/Calmodulin-dependent protein kinase I (CAMKI), creatine transporter (CRTR), CDM protein (CDM) adramoleukogusekontus medelen
4042	13846	23724	2.25	1.0E-10	AB031069.1	ĽΝ	Homo series BCCV1 mBMA 4
679	13981		2.06	1.0E-10		NT	Human pregnancy appetiting a property and 1 (CDA) That a
5145	15012		0.93	1.0E-10 >	X87344.1	Z	H. sepiens DMA, DMB, HLA-Z1, IPP2, LMP2, TAP1, LMP7, TAP2, DOB, DQB2 and RING8, 9, 13 and 14 genes
7859	17709		4.59	1.0F-10.4	0 A DR 1868 1		
8286	18165	28408	3.4	_	1	FST HIMAN	on School of Street School of School of Street School of School of Street School of Street School of S
9037	15012		54.1	1.0E-10		\neg	H. Sepiens D.M.A. DMB, HLA-Z1, IPP2, LMP2, TAP1, LMP7, TAP2, DOB, DQB2 and RINGB, 9, 13 and 14
9084	18859		1.3	1.0E-10 A	5.1	T HUMAN	288610 C Sparse toolle NUT Long
					1	1	CONTROL OCER SEALS TO TO SAPIENS CONA CIONE IMAGE: 729211 F.

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Table 4
Single Exon Probes Expressed in Heart

					•		
Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
261	10226	20042	1.2	8 11-30'6	BE145600.1	EST_HUMAN	IL2-HT0203-291099-016-c08 HT0203 Homo sapiens cDNA
2037			5.43		9.0E-11 AL134395.1	EST_HUMAN	DKFZp547D225_r1 547 (synonym: hfbr1) Homjo sapiens cDNA clone DKFZp547D225 5'
2057			5,43		9.0E-11 AL134395.1	EST_HUMAN	DKFZp547D225_r1 547 (synonym: hfbr1) Homo sapiens cDNA clone DKFZp547D225 5'
3337			2.25		9.0E-11 AL134395.1	EST_HUMAN	DKFZp547D225_r1 547 (synonym: hfbr1) Homo sapiens cDNA clone DKFZp547D225 5'
3337		23065	2.25		İ	EST_HUMAN	DKFZp547D225_r1 547 (synonym: hfbr1) Homo sapiens cDNA clone DKFZp547D225 5'
4400			1.09			EST_HUMAN	ae78f01.s1 Stratagene schizo brain S11 Homo saplens cDNA clone IMAGE:970297 3'
5421	15342		4	9.0E-11	1.	EST_HUMAN	RC6-BT0627-140200-011-E06 BT0827 Homo sapiens cDNA
8410	18062	25312	2.69		9.0E-11 C16635.1	EST_HUMAN	C16635 Clontech human aorta polyA+ mRNA (#6572) Homo sapiens cDNA clone GEN-506B08 5'
	l						yn63111.s1 Soares adult brain N2b5HB55Y Homo sapiens cDNA clone IMAGE:172173 3' similar to contains
3078	13005		9.97	8.0E-11	H19971.1	EST_HUMAN	L1 repetitive element ;
3881		23580	0.83	8.0E-11	8.0E-11 AI478617.1	EST_HUMAN	tm54c09.x1 NCI_CGAP_Kid11 Homo saplens ¿DNA clone IMAGE:2161936 3'
3957			4.03	8.0E-11	N23712.1	EST_HUMAN	yw46e06.s1 Welzmann Olfactory Epithelium Homo sapiens cDNA clone IMAGE:255298 31
1431	11336	21202	2.11	7.0E-11	7.0E-11 AA330642.1	EST_HUMAN	EST34392 Embryo, 6 week I Homo sapiens cONA 5' end
6949	16827		2.55	7.0E-11 A	AF163864.1	NT	Homo sapiens SNCA isoform (SNCA) gene, complete cds, alternatively spliced
							RETROVIRUS-RELATED POLYPROTEIN [CONTAINS: REVERSE TRANSCRIPTASE;
7873	17723	•	1.22	7.0E-11		SWISSPROT	ENDONUCLEASE]
9545			1.31		3.1	EST_HUMAN	AV701656 ADB Homo sapiens cDNA clone ADBABC09 5'
408	1		5.16		8.0E-11 M55270.1	NT	Human matrix Gla protein (MGP) gene, complete cds
408			5.16		6.0E-11 M55270.1	NT	Human matrix Gla protein (MGP) gene, complete cds
6259		26568	3.58		6.0E-11 P08547	SWISSPROT	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG
6871	Ĺ		6.49			EST_HUMAN	AV727859 HTC Homo sapiens cDNA clone HTCASC08 5'
11		19788	0.92		5.0E-11 AL163283.2	NT	Homo sapiens chromosome 21 segment HS210083
3320	2668	19788	1.48		5.0E-11 AL163283.2	LN	Homo saplens chromosome 21 segment HS21C083
4135			1.49		5.0E-11 P48034	SWISSPROT	ALDEHYDE OXIDASE
2926		25954	1.69		AL163213.2	LN.	Homo sapiens chromosome 21 segment HS21C013
6446			14.59		11416799	TN	Homo sapiens protocadherin beta 3 (PCDHB3) mRNA
8976	18780		1.79	5.0E-11	5.0E-11 AJ289880.1	LN	Homo sapiens KIAA0851 gene (partial), XT3 gene and LZTFL1 gene
1380	11285		1.4		4.0E-11 AA436042.1	EST_HUMAN	zu01b12.r1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:730559 5'
2760	12622	22514	8.45		4.0E-11 BE885900.1	EST_HUMAN	601507531F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3909295 5'
4518	14411	24196	1.37	4.0E-11	4.0E-11 D44668.1	EST_HUMAN	HUMSUPY069 Human brain cDNA Homo sepiens cDNA clone 069
5897	15803	25927	2.94	4.0E-11	4.0E-11 P20095	SWISSPROT	PRE-MRNA SPLICING FACTOR RNA HELICASE PRP2
6362	16225		3.66		AF224669.1	TN	Homo sapiens mannosidase, beta A, lysosomal(MANBA) gene, and ubiquitin-conjugating enzyme E2D 3 (UBE2D3) genes, complete cds
7405	17272		1.59		4.0E-11 BE149425.1	EST_HUMAN	RC1-HT0256-210100-013-f08 HT0256 Homo saplens cDNA

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Top Hit Descriptor	Home sapiens SH3-domain binding protein 1 (SH3BP1), mRNA	Mus musculus expressed in non-metastatic cells, 2, protein (NM23B) (Nme2), mRNA	EST180120 Liver, hepatocellular carcinoma Homo sapiens cDNA 5' end	qf36c04.x1 Soares_tests_NHT Homo saplens cDNA clone IMACE:1752102.3' similar to contains MER10.t3 MER10 repetitive element:	Τ	19843912.r1 Soares Infant brain 1NIB Homo sapiens cDNA clone IMAGE:35144 5	Gellus gallus rho-globin, beta-H globin, beta-A globin, epsilon-globin, and olfactory receptor-like protein COR3 beta (COR3 beta) genes, complete cds	Gallus gallus rho-globin, beta-H globin, beta-A globin, epsilon-globin, and olfactory receptor-like protein COR3'beta (COR3'beta) genes, complete cds	qc51c10.x1 Soares_pregnant_uterus_NbHPU Homo sapiens cDNA clone IMAGE:1713138 3' similar to gb:L02932 PEROXISOME PROLIFERATOR ACTIVATED RECEPTOR ALPHA (HUMAN);contains L1.t1	L1 repetitive element;	Human endogenous retrovirus HERV-P-T47D	RETROVIRUS-RELATED GAG POLYPROTEIN (VERSION 1)	m54c09.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2161936 3	Homo sapiens FRA3B common fragile region, diadenosine triphosphate hydrolase (FHIT) gene, exon 5	Homo sapiens chromosome 21 segment HS21C027	QV2-BT0258-261099-014-a01 BT0258 Homo sapiens cDNA	EST178226 Colon carcinoma (HCC) cell line Homo sapiens cDNA 5' end similar to similar to alpha-2-macroglobulin	nc83h05.r1 NCI_CGAP_GC1 Homo sapiens cDNA clone IMAGE:797433 5' similar to SW:PR16_YEAST P15938 PRE-MRNA SPLICING FACTOR RNA HELICASE PRP16.;	Home sapiens chromosome 9 duplication of the Toeli receptor beta focus and tropsinocen gene families	OLFACTORY RECEPTOR 511 (OLFACTORY RECEPTOR-LIKE PROTEIN OLF1)	zk27g02.s1 Soares_pregnant_uterus_NbHPU Homo sapiens cDNA clone IMAGE:471794 3'	zk27g02.s1 Soares_pregnant_uterus_NbHPU Homo sapiens cDNA clone IMAGE:471794 3'	277e03.s1 Soares_fetal_liver_spleen_1NFLS_\$1 Homo sapiens cDNA clone IMAGE:460924.3'	RC0-CN0027-210100-011-c01 CN0027 Homo sapiens cDNA	CM2-TN0140-070900-372-g01 TN0140 Homo sapiens cDNA	Homo sapiens mRNA for KIAA0027 protein, partial cds
Top Hit Database Source	NT	TN	EST_HUMAN	FST HUMAN	EST HUMAN	EST HUMAN	FZ	FZ		EST_HUMAN	TN	SWISSPROT	EST_HUMAN	TN	TN	EST_HUMAN	EST_HUMAN	EST_HUMAN	LN	SWISSPROT	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	N
Top Hit Acession No.	11545732 NT	6679077 NT	AA309248.1	A1150502 1	R24807.1	R24807.1	L17432.1	L17432.1		AI128371.1	AF087913.1	P10263	AI478617.1	AF020503.1	AL163227.2	BE062558.1	AA307331.1	AA581028.1	AF029308.1	Q13606	AA035369.1	AA035369.1	AA704195.1	AW842143.1	BF377859.1	D25217.2
Most Similar (Top) Hit BLAST E Value	4.0E-11	3.0E-11	3.0E-11	. 2.0E-11	_		2.0E-11	2.0E-11			20E-11		2.0E-11		2.0E-11	2.0E-11	2.0E-11	2.0E-11	2.0E-11	_			2.0E-11		2.0E-11	2.0E-11
Expression Signal	1.93	2.18	1.45	1.68	3.64	3.64	3.91	3.91		1.04	1,11	4.23	0.87	0.85	76.0	5.48	1.32	1.83	1.43	4.61	2.07	2.07	1.29	2.25	1.98	1.89
ORF SEQ ID NO:	25252			20718				21360				22889					24804	25828		27989		28592			25356	
Exan SEQ ID NO:	19190		14079	10869		L	11500	11500			12599		13205		14395	14732	14836	15715	17194	\mathbf{L}_{-}	18330	18330	19679		Ιí	19103
Probe SEQ ID NO:	9611	1478	4178	944	1168	1168	1596	1598		1600	2737	3160	3284	3452	4501	4851	4961	5810	7318	7900	8457	8457	9160	9192	9216	9477

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.		Top Hit Descriptor
9627	19201		2.38	-	P08547	SWISSPROT	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG
8930	19401		2.38	2.0E-11	11417966 NT	TN	Homo sapiens SEC14 (S. cerevisiae)-like 2 (SEC14L2), mRNA
980	10594	20412	1.24		AJ131016.1	TN	Homo saplens SCL gene locus
1199			2.58	1.0E-11	AL163279.2	LN	Homo saplens chromosome 21 segment HS21C079
1483			1.94			NT	Homo sapiens PRO3078 mRNA, complete cds
. 2079		21862	3.12			NT	Homo saplens homogentisate 1,2-dioxygenase gene, complete cds
3454			1.32		BE004315.1	EST_HUMAN	CMO-BN0105-170300-292-d12 BN0105 Homo sapiens cDNA
5269		24966	14.34	1.0E-11	AL163247.2	NT	Homo sapiens chromosome 21 segment HS21C047
6773	il		3.25	1.0E-11	4885546 NT		Homo sapiens PHD finger protein 2 (PHF2) miRNA
6869	16868		5.41		R13174.1	EST_HUMAN	y73d08.r1 Soares Infant brain 1NIB Homo sapiens cDNA clone IMAGE;28168 5'
7232	17109		1.32		BF365119.1	EST_HUMAN	QV4-NN1149-250900-423-803 NN1149 Homo sapiens cDNA
7232	17109	27301	1.32	1.0E-11	BF365119.1		QV4-NN1149-250900-423-a03 NN1149 Home saplens cDNA
8607			1.9		BF680078.1	1	602154807F1 NIH_MGC_83 Homo saplens cDNA clone IMAGE:4295977 5'
9702			1.29	1.0E-11	Z20377.1		HSAAACADH P, Human foctal Brain Whole tissue Homo expiens cDNA
2922	12849	22849	0.82	9.0E-12	P20742	SWISSPROT	PREGNANCY ZONE PROTEIN PRECURSOR
7641		27712	1.22	9.0E-12		NT	Homo sapiens chromosome 21 segment HS21C100
7641		27713	1.22			NT	Homo sapiens chromosome 21 segment HS21C100
9270	18974		3.57	8.0E-12	16.1		Homo sapiens Xq pseudoautosomal region; segment 2/2
4562		24241	2.75	7.0E-12	005904		34 KD SPICULE MATRIX PROTEIN PRECURSOR (LSM34)
8658	3 18545		11.23	7.0E-12	AA704735.1		423g01.s1 Scares_fettal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:4511523'
3500	13417		0.92	6.0E-12	AV730554.1	EST_HUMAN	AV730554 HTF Homo sapiens cDNA done HTFAW F06 5'
			F		. 072000		nz88f11.s1 NCI_CGAP_CCB1 Homo sapiens cDNA clone IMAGE:1302573 3' similar to contains Alu
7204	17076	27282	60.7	9.0E-12		FIGURE FIN	Mycone caxallis mysein beam chain EM34 (EM34) mRNA complete cde
	1					7	od10d11 s1 NCI CGAP GCB1 Homo seniens CDNA clone IMAGE:1367588 similar to contains MER29.12
7463	17262		1.86	8.0E-12	AA847898.1	EST_HUMAN	MER29 repetitive element;
1027	7 10945		2.88	5.0E-12	T06573.1	EST_HUMAN	EST04462 Fetal brain, Stratagene (cat#936206) Homo sapiens cDNA clone HFBDV33
3344		02002	1.18	5.0E-12	BE047779.1	EST_HUMAN	tz42b05.y1 NCI_CGAP_Brn52 Homo saplens cDNA clone IMAGE:2291217 5
3666			5.83	5.0E-12	AJ271736.1	NT	Homo saplens Xq pseudoautosomal region; segment 2/2
5171			0.84	5.0E-12	AA720661.1	T_HUMAN	nw24b11.s1 NCI_CGAP_GCB0 Homo sapiens cDNA clone IMAGE:1241373 3'
5667			4.75	5.0E-12	AL163278.2	NT	Homo sapiens chromosome 21 segment HS210078
2999			4.75	5.0E-12		NT	Homo sapiens chromosome 21 segment HS21C078
5909	1	25940	9.12	5.0E-12	5.0E-12 AW974760.1	T_HUMAN	EST386850 MAGE resequences, MAGN Homo sapiens cDNA
7268	17145	1	2.15	5.0E-12		Ę	Homo saplens Xq pseudoautosomal region; segment 1/2

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Single Exon Probes Expressed in Heart

1756 17745 2002 3.45 4.0E-12/A-V1002031 EST HUMAN 17414 18 Sares, feat liver, galement ARX 19 Sares 17745 24 Sares 17415 24 Sares 17415 24 Sares 17415 24 Sares 17415 24 Sares 17415 24 Sares 17415 24 Sares 17415 24 Sares 17415 24 Sares 17415 24 Sares 17415 24 Sares	Probe SEQ ID NO:	SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
10211 20027 3.42 4.0E-12 AA700326.1 EST_HUMAN 10211 20027 3.55 4.0E-12 AA700326.1 EST_HUMAN 14413 24198 0.86 4.0E-12 AF108904.1 EST_HUMAN 16676 2.89 4.0E-12 AF108907.1 NT 18305 28561 3.51 4.0E-12 AF108907.1 NT 10536 20346 3.81 3.0E-12 AW341683.1 EST_HUMAN 10538 20346 3.81 3.0E-12 AW341683.1 EST_HUMAN 17839 28187 3.08 3.0E-12 AW341683.1 EST_HUMAN 13828 23704 1.04 2.0E-12 AW341683.1 NT 13828 23704 1.04 2.0E-12 AW341683.1 NT 13828 23704 1.04 2.0E-12 AW341683.1 NT 14221 1.03 2.0E-12 AW341683.1 NT 14828 2304 1.04 2.0E-12 AW341683.1 NT 14829 23704 1.04 2.0E-12 BE083509.1 NT 14888 244	7895	L.		4.67	6.0E-12	AL163303.2	TN	Homo saplens chromosome 21 segment HS2IC103
10211 20027 3.56 4.0E-12 AA700326.1 EST_HUMAN 14413 24198 0.86 4.0E-12 AF109907.1 NT 16676 2.89 4.0E-12 AF109907.1 NT 18305 28561 3.51 4.0E-12 AF109907.1 NT 10536 280346 3.81 3.0E-12 AW341683.1 EST_HUMAN 10538 20346 3.81 3.0E-12 U78027.1 NT 17838 228187 3.08 3.0E-12 U37672.1 NT 17838 23148 3.08 3.0E-12 U37672.1 NT 13828 23704 1.04 2.0E-12 U37672.1 NT 14221 1.37672.1 NT NT 13828 23704 1.04 2.0E-12 U37672.1 NT 14221 1.37672.1 NT NT 14221 1.03 2.0E-12 U37672.1 NT 14221 0.78 2.0E-12 U384.1 NT 14828 24473 0.78 2.0E-12 U7864.1 NT 14839	244	Ш		3.42	4.0E-12		EST_HUMAN	474g11.s1 Soares_fetal_liver_spleen_1NFLSi_S1 Homo saplens cDNA clone IMAGE:460676 3'
14413 24198 0.86 4.0E-12 AI6B9B84.1 EST_HUMAN 16676 2.89 4.0E-12 AI2B907.1 NT 18305 28561 3.51 4.0E-12 AI2B907.1 NT 10536 20346 3.81 3.0E-12 AW341683.1 EST_HUMAN 17838 28187 3.0E-12 AW341683.1 EST_HUMAN 17838 28187 3.0E-12 U37672.1 NT 17838 28187 3.0E-12 U37672.1 NT 13928 23704 1.04 2.0E-12 U37672.1 NT 14828 23473 1.04 2.0E-12 U37672.1 NT 14828 23474 0.78 2.0E-12 U37672.1 NT 14828 23474 0.78 2.0E-12 U37672.1 NT 14829 23704 1.04 2.0E-12 U37672.1 NT 14820 2375 1.04 2.0E-12 U376306 SWISSPROT 15804 2.22 2.0E-12 U376306 SWISSPROT 1681 2.07 2.0E-12 U376306 SWISSPROT <t< td=""><td>245</td><td></td><td>L</td><td>3.55</td><td>4.0E-12</td><td>AA700326.1</td><td>EST_HUMAN</td><td>474g11.s1 Soares_fetal_liver_spleen_1NFLSLS1 Homo sapiens cDNA clon⊌ IMAGE:460676 3'</td></t<>	245		L	3.55	4.0E-12	AA700326.1	EST_HUMAN	474g11.s1 Soares_fetal_liver_spleen_1NFLSLS1 Homo sapiens cDNA clon⊌ IMAGE:460676 3'
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18305 28561 3.51 4.0E-12 AJ29043.1 NT 16316 20346 3.81 3.0E-12 AJ29043.1 NT 10536 20346 3.81 3.0E-12 AW341683.1 EST_HUMAN 17838 28187 3.08 3.0E-12 U37672.1 NT 17838 23143 1.03 2.0E-12 U37672.1 NT 13928 23704 1.04 2.0E-12 U37672.1 NT 14221 1.03 2.0E-12 U37672.1 NT 14829 24473 0.78 2.0E-12 U37689.1 EST_HUMAN 1584 2.27 3.34 2.0E-12 AW371857.1 EST_HUMAN 16124 26277 3.34 2.0E-12 AW371857.1 EST_HUMAN 16376 26652 2.07 2.0E-12 AW871857.1 EST_HUMAN 16376 26652 2.07 2.0E-12 AW871857.1 EST_HUMAN 16377 1.52 2.0E-12 AW871857.1 EST_HUMAN 16376 16941 1.52 2.0E-12 AW871857.1 EST_HUMAN 1838 2478 1.0E-12 AW871857.1 EST_HUMAN 1848 2478 1.0E-12 AF165283.2 NT 1858 1.0E-12 AF165283.2 NT 1859 2.03 1.0E-12 AF060991.1 NT 1789 22752 1.16 1.0E-12 AF009991.1 NT 1789 28752 1.16 1.0E-12 AF009991.1 NT 1780 28752 1.16 1.0E-12 AF00991.1 NT 1780 28752 1.16 1.0E-12 AF00909	1000	٠.				A 17,00004 4	E.V	Homo sapiens S164 gene, partial cds; PS1 and hypothetical protein genes, complete cds; and S171 gene, $\hat{\sigma}$
19131 1.9 4.0E-12 U78027.1 NT 10536 20346 3.81 3.0E-12 AW341683.1 EST_HUMAN 17838 28187 3.08 3.0E-12 U37672.1 NT 17838 23143 1.03 2.0E-12 U37672.1 NT 13828 23704 1.04 2.0E-12 J01884.1 NT 14829 2477 0.78 2.0E-12 J01884.1 NT 14839 2477 0.78 2.0E-12 J01884.1 NT 15804 2.22 2.06-12 AW371857.1 EST_HUMAN 16376 2652 2.07 2.0E-12 AW371857.1 EST_HUMAN 16376 2652 2.07 2.0E-12 AW371857.1 EST_HUMAN 16376 2652 2.07 2.0E-12 AW371857.1 EST_HUMAN 16376 2652 2.07 2.0E-12 AW371857.1 EST_HUMAN 16376 2652 2.07 2.0E-12 AW371857.1 EST_HUMAN 16376 2652 2.07 2.0E-12 AW371857.1 EST_HUMAN 16376 2652 2.07 2.0E-12 AW371857.1 EST_HUMAN 16376 2652 2.07 2.0E-12 AW371857.1 EST_HUMAN 16376 2652 2.07 2.0E-12 AW371857.1 EST_HUMAN 16376 2652 2.07 2.0E-12 AW371857.1 EST_HUMAN 16376 16914 1.52 2.0E-12 AF163283.2 NT 16085 16914 1.62 2.03 1.0E-12 AF163283.2 NT 17890 2.03 1.0E-12 AF163283.1 NT	8431	L		3.51		AJ229043.1	LZ	Homo saplens 959 kb contig between AML1 and CBR1 on chromosome 21q22, Caginent 3/3
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13338 23143 1.03 2.0E-12 675495 NT 13928 23704 1.04 2.0E-12 J01884.1 NT 13928 23704 1.04 2.0E-12 J01884.1 NT 14221 1.04 2.0E-12 J01884.1 NT 1488 2.473 0.78 2.0E-12 BE063509.1 EST_HUMAN 1488 2.474 0.78 2.0E-12 AW917857.1 EST_HUMAN 15804 2.22 2.0E-12 AW917857.1 EST_HUMAN 16376 2.62 2.0E-12 AW917857.1 EST_HUMAN 17345 1.68 2.0E-12 AF196964.1 NT 17346 1.71 2.0E-12 AF196964.1 NT 1837 8.13 2.0E-12 AF196864.1 NT 19041 1.77 2.0E-12 AF196864.1 NT 19041 1.52 2.0E-12 AF196864.1 NT 19041 1.52 2.0E-12 AF196864.1 NT </td <td>8047</td> <td>L</td> <td></td> <td></td> <td>3.0E-12</td> <td></td> <td>LN</td> <td>Human prostate specific antigen gene, 5' flanking region</td>	8047	L			3.0E-12		LN	Human prostate specific antigen gene, 5' flanking region
13928 23704 1.04 2.0E-12 JO1884.1 NT 13828 23705 1.04 2.0E-12 JO1884.1 NT 14221 1.04 2.0E-12 JO1884.1 NT 14828 2.4473 0.78 2.0E-12 JO70306 SWISSPROT 14888 2.4474 0.78 2.0E-12 JW871857.1 EST_HUMAN 15804 2.22 2.0E-12 JW871857.1 EST_HUMAN 16376 2.05 2.0E-12 JW871857.1 EST_HUMAN 17345 3.34 2.0E-12 JAT18684.1 NT 17345 8.13 2.0E-12 JAT18684.1 NT 18976 1.77 2.0E-12 JAT18684.1 NT 18976 1.77 2.0E-12 JAT18288.1 NT 19041 1.52 2.0E-12 JAT18248 NT NT 10085 19914 1.82 1.0E-12 JAT18248 NT EST_HUMAN 11889 2.03 1.0E-12 JAR1728.1 EST_HUMAN 12860 2.03 1.0E-12 JAR2728.1 EST_HUMAN 12860 2.05 1.0E-12 JAR27728.1 <td>3421</td> <td>L</td> <td></td> <td></td> <td>2.0E-12</td> <td>6754495</td> <td>TN</td> <td>Mus musculus keratin-associated protein 6.2 (Krtap6-2), mRNA</td>	3421	L			2.0E-12	6754495	TN	Mus musculus keratin-associated protein 6.2 (Krtap6-2), mRNA
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1468B 24473 0.78 20E-12 O70306 SWISSPROT 1468B 24474 0.76 20E-12 O70306 SWISSPROT 15804 2.22 20E-12 AW971857.1 EST_HUMAN 16124 28277 3.34 20E-12 AW971857.1 EST_HUMAN 1637B 2652 2.07 2.0E-12 AF18684.1 EST_HUMAN 17345 1,68 2.0E-12 AF186864.1 NT 18916 1,71 2.0E-12 AF186864.1 EST_HUMAN 19041 1,52 2.0E-12 AF182880.1 EST_HUMAN 10085 18914 1,52 2.0E-12 AF18248.1 EST_HUMAN 11839 2.03 1.0E-12 AF1824.1 EST_HUMAN 1839 2.03 1.0E-12 AF000991.1 EST_HUMAN	4324	L_		1.8			EST_HUMAN	GM0-BT0281-031199-087-a03 BT0281 Homo saplens cDNA
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16376 26552 2.07 2.0E-12 AF196864.1 11422229 NT 17345 1,68 2.0E-12 AF196864.1 NT 17587 8.13 2.0E-12 BE165980.1 EST_HUMAN 18916 1,71 2.0E-12 AL163283.2 NT 10041 1,52 2.0E-12 AL163283.2 NT 10095 19914 1.82 1.0E-12 AW627674.1 EST_HUMAN 12950 22762 1,16 1.0E-12 AF000991.1 NT	8228	L				T08169.1	EST_HUMAN	EST06060 Infant Brain, Bento Scares Homo sapiens cDNA clone HIBBA13 5' end
17345 1.68 2.0E-12 AF198864.1 NT 17587 8.13 2.0E-12 BE165980.1 EST_HUMAN 18916 1.71 2.0E-12 AL163283.2 NT 10041 1.52 2.0E-12 AL163283.2 NT 10095 19914 1.82 1.0E-12 AW627674.1 EST_HUMAN 11839 2.03 1.0E-12 AR21726.1 EST_HUMAN 12960 22762 1.16 1.0E-12 AF000991.1 NT	8516	L				11422228	LN	Homo sapiens Ac-like transposable element (ALTE), mRNA
17587 8.13 2.0E-12 BE165980.1 EST_HUMAN 18916 1.71 2.0E-12 AL163283.2 NT 19041 1.52 2.0E-12 AL163283.2 NT 10095 18914 1.82 1.0E-12 AW627674.1 EST_HUMAN 11839 2.03 1.0E-12 AR71726.1 EST_HUMAN 12960 22752 1.16 1.0E-12 AF000991.1 NT	7367	<u> </u>				AF196864.1	IN	Hamo sapiens putative BPES syndrome breakpoint region protein gene, complete cds
18916 1.71 2.0E-12 AL163283.2 NT 19041 1.52 2.0E-12 AL163248 NT 10095 18914 1.82 1.0E-12 AW627674.1 EST_HUMAN 11839 2.03 1.0E-12 AR71726.1 EST_HUMAN 12960 22762 1.16 1.0E-12 AF000991.1 NT	737	L		8.13			EST_HUMAN	MR3-HT0487-150200-113-g01 HT0487 Homo saplens cDNA
19041 1.52 2.0E-12 11418248 NT 10055 18914 1.82 1.0E-12 AW627674.1 EST_HUMAN 11839 22762 1.16 1.0E-12 AR500991.1 NT	9175	L		1.71	2.0E-12		¥	Hamo saplens chromosome 21 segment HS21C083
10095 18914 1.82 1.0E-12 AW627674.1 EST_HUMAN 11839 22762 1.16 1.0E-12 AF000991.1 NT	837			1.52	2.0E-12		Į,	Homo sapiens sulfotransferase-related protein (SULTX3), mRNA
10096 18914 1.82 1.0E-12 AW627674.1 EST_HUMAN 11839 22762 1.16 1.0E-12 AF000991.1 NT								hh90a09.x1 NCI_CGAP_GU1 Homo saplens cDNA clone IMAGE:2970040 3' similar to contains MER18.t1
11839 2.03 1.0E-12 AI871726.1 EST_HUMAN 12960 22762 1.16 1.0E-12 AF000991.1 NT	117					AW627674.1	EST_HUMAN	MER18 repetitive element ;
12960 22762 1.16 1.0E-12 AF000891.1 NT	194	1		2.03		AI871726.1	EST_HUMAN	wm51f07.x1 NCI_CGAP_Ut2 Homo sapiens clDNA clone IMAGE:2439493 3' straitar to contains L1.b3 L1 repetitive element ;
	3032	1 1				AF000991.1	NT.	Homo sapiens testis-specific Testis Transcript Y 2 (TTY2) mRNA, partial cds

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Table 4
Single Exon Probes Fynnam

Protect Expr. Opp. Str. Expression Trop-Hill Tro-Hill Above Stylings Tro-Hill Tro-Hill Above Stylings Tro-Hill Tro-Hill Above Stylings Tro-Hill Tro-Hill Above Stylings Tro-Hill Tro-Hill Above Stylings Tro-Hill Tro-Hill Above Stylings Tro-Hill Tro-Hill Above Stylings
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18860 29118 3.36 1.0E.12 AWGGGGG
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Single Exon Probes Expressed in Heart

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Single Exon Probes Expressed in Heart	Top Hit Descriptor	WROTH A Secret Plane in 1911	zitzen. 1 Otares pacerila Nazir Fromo sapiens CDNA clone IMAGE:145759 5' zitzetz.st Soares_testis_NHT Homo sapiens cDNA clone IMAGE:728350 3' similar to contains Alvi	l'opeduve element; confains element MER22 repetitive element;	MYOSIN LIGHT CHAIN KINASE, SKELETAL MUSCLE (MLCK)	PM2-H 10224-221099-001-e11 HT0224 Homo sapiens cDNA	Homo sapiens glypican 3 (GPC3) gene, partial cds and flanking repeat regions	rws-r1 0520-230200-002-c08 H10520 Homo septens cDNA	y63955 f complex resilience of similar to PIR: A32995 f complex resilience of similar to PIR: A32995 f complex resilience of similar to PIR: A32995	qn32d05x1 NCI_CGAP_Kld5 Homo saplens cDNA clone IMAGE:1899945 3' similar to contains Alu	repeuve dement;	278910.51 Soares (estis_NHT Homo sapiens cDNA clone IMAGE:728514.3'	Homo seplens X-linked anhidrolite ectodermal dysplasia protein gene (EDA), exxn. 2 and flanking reneat	Sholler	Avoigue, r.1 Soares_testis_NHT Homo saplens cDNA clone IMAGE:781406 5	namo sapiens Aq pseudoautosomal region; segment 2/2	CM2 Erzess discussione 21 segment HS21C010	ONIS-F1010U-14U/00-242-h08 FT0100 Homo sapiens cDNA	Homo sapiens X28 region near ALD locus containing dual specificity phosphatase 9 (DUSP9), ribosomal protein L18a (RPL18a), Ca2+/Calmodulin-dependent protein kinasa (CAMM)	CDM protein (CDM), adrenoleukodystrophy protein >	HA0536 Human fetal liver cDNA library Homo sapiens cDNA	CMC-B10281-031199-087-a03 BT0281 Homo (appiens cDNA	Flomo septens chromosome 21 segment HS21(5048	protein L18a (RPL18a). Ca2+/Calmodulin-dependent protein kinase I (CAMKI), creatine transporter (CRTR)	Omio rario filtrahlasi manda ƙasar Santa Angela Santa Rario filtrahlasi manda ƙasar Santa	Dance of the control	nomo sabiens UNA polymerase delta small subiunit (POLD2) gene, exons 1 through 11 and complete cds nab76t05.x1 Soares NSF F8 9W OT PA B S1 Library	STATE OF THE STATE
gle Exon Pro	Top Hit Database Source	EST HUMAN		EST HOMAN	SWISSPROI	ESI HUMAN	COT LIMANN	LA LA	EST HUMAN	CT 11 1444.1	EST LINEAR	EST HIMAN	±N		Т		T 11 184581	Т	NCA	1011111111	T	NAW OF THE	\int				T_HUMAN	ı
ilo I	Top Hit Acession No.	3 R78338.1	3 44435779 4	T	27.7	7	T					T		T	T	T	T	T		052111.2 NT		T	10170017	U62111.2 NT				
	Most Similar (Top) Hit BLAST E Value	5.0E-13	5.05-13	50E-13	4.0F-13	4 OF-13	4.0E-13	4.0E-13	4.0E-13	4.0F-13	4.0F-13/	4.0E-13/	3.0E-13.4	3.0E-13.A	3.0E-13 A	3.0E-13.A	3.0E-13/B	3.0E-13 A	0	3.0E-13.0	3 0F-13 R	3 0F 43 A	2	2.0E-13 U	2.0E-13 U;		2.0E-13 BF	
	Expression Signal	1.05	1.45	2.75	2.96	1.58	4.86	1.92	1.52	4.57	1.83	1.83	3.8	1.37	1.25	2.63	4.23	2.86	c u	4.03	3.66	2.6		2.77	1.31	4.71	1.08	
	ORF SEQ ID NO:			28367			25404			27831	28645	28646			22099		22387	-	26680		28531	28980	-	19938	20024	21005	22966	
	(I)	13201	13271	18115	11721	12288	L	16142	16400	17607	18379	18379	10144	10774	12200	12306	12497	13075	16481	17955	18279	18688		10118	10207	11156	13167	
	Probe SEQ ID NO:	3280	3351	8234	1824	2411	5430	6278	6542	77.57	8507	8507	173	847	2319	2429	2629	3150		8064	8403	8876	-	144	82	1249	3244	

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Single Exon Probes Expressed in Heart

Top Hit Descriptor	Homo sapiens chromosome 21 segment HS210078	CELL SURFACE GLYCOPROTEIN 1 PRECURSOR (OUTER LAYER PROTEIN B) (S-LAYER PROTEIN 1)	Human PFKL gene for liver-type 6-phosphofruciokinase (EC 2.7.1.11) exon 2	Homo sapiens mab-21 (C. elegans)-like 1 (MAE21L1) mRNA	CM0-NN0001-100300-274-e11 NN0001 Homo sapiens cDNA	FGF-1=fibroblast growth factor 1 [human, kidney, Genomic, 342 nt, segment 2 of 2]	Homo sapiens LGMD2B gene	H.sapiens DMA, DMB, HLA-Z1, IPP2, LMP2, TAP1, LMP7, TAP2, DOB, DQB2 and RING8, 9, 13 and 14 dense.	pwo/and st NCI (33AP 3CR0 Homo seniens InDNA clone IMAGE 1241138 3' similar to contains THB t3	THR repetitive element;	602038009F1 NCI_CGAP_Brn64 Homo sapiens cDNA done IMAGE:4185866 5	745e10.x1 Scares_NSF_F8_9W_OT_PA_P_S1 Homo sapiens cDNA clone IMAGE:3524443.3' similar to	contains MER29.b2 MER29 repetitive element ;,	AV715377 DCB Homo sapiens cDNA clone DQBAIE03 5	Homo sapiens Xq pseudoautosomal region; segment 1/2	aj24001.s1 Soares_testis_NHT Homo sapiens dDNA clone 1391232 3' similar to contains MER19.t1 MER19 repetitive element;	al 24c01.s1 Soares, testis NHT Homo sapiens cDNA clone 1391232.3° cimilar to contains MER19.11 MER19	repetitive element;	RC4-CT0322-080100-013-d09 CT0322 Homo saplens cDNA	Homo sapiens TFF gene cluster for frefoil factor, complete cds	xo54h05.x1 NCI_CGAP_Ut1 Homo sapiens cDNA clone IMAGE:2707833 3'	aj24c01.s1 Soares_testis_NHT Homo sapiens cDNA clone 1391232 3' similar to contains MER19.11 MER19	repetitive element ;	Human DNA, SINE repetitive element	Saguinus oedipus gene for seminal vesicle secreted protein semenogelin i	hz71c09.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3213424 3'	y72e03.r1 Soares placenta Nb2HP Homo sapiens cDNA clone IMAGE:1447963'	H.sapiens DNA for endogenous retroviral like element	zq17c10.s1 Stratagene fetal retina 937202 Horrio sapiens cDNA clone IMAGE:629970 3'	QV2-BT0258-261099-014-a01 BT0258 Homo sapiens cDNA
Top Hit Database Source	LN	SWISSPROT	N		EST_HUMAN	TN	IN	F		EST HUMAN			HUMAN	T_HUMAN	LN	EST HUMAN	1	EST_HUMAN	EST_HUMAN	LN.	EST_HUMAN		EST_HUMAN	LN	FZ		T_HUMAN		П	EST HUMAN
Top Hit Acession No.	3 AL163278.2	Q06852	2.0E-13 X16912.1	5031896 NT	AW892155.1	1.0E-13 S74129.1	3 AJ007973.1	4 0E-13 X87344 1		1.0E-13 AA720574.1	3 BF340987.1		1.0E-13 BF108755.1	3 AV715377.1	1.0E-13 AJ271735.1	4 AA781159.1		9.0E-14 AA781159.1	9.0E-14 AW861577.1	9.0E-14 AB038162.1	4 AW513298.1		4 AA781159.1	4 D14547.1	4 AJ002153.1	4 BE468263.1	4 R76269.1	4 X89211.1	4 AA219316.1	4 BE062558.1
Most Similar (Top) Hit BLAST E Value	2.0E-13	2.0E-13	2.0E-13	2.0E-13	2.0E-13	1.0E-13	1.0E-13	4 OF.43		1.0E-13	1.0E-13		1.0E-13	1.0E-13	1.0E-13	9.0E-14		9.0E-14	9.0E-14	9.0E-14	9.0E-14		9.0E-14	9.0E-14	9.0E-14	8.0E-14	8.0E-14	8.0E-14	8.0E-14	8.0E-14
Expression Signal	1.72	3.87	6.32	3.97	7.42	1.37	4.39	1 97		2.16	1.48		13.83	1.62	1.6	2.92		2.85	4.04	4.62	3.74		0.98	5.22	1.93	1.67	2.77	69.69	3.49	4.39
ORF SEQ ID NO:			26209			20072	20848				24169		28855			20105		20108	-		22790			23428	24325			26637	27510	
Exen SEQ ID NO:	13923	15629	16061	Į	1	. 10252	10798	11220	_	11869	Į.		_	.)	19255	10289		10290	ı	l	13000	1	10289	13640				1	1 1	18588
Probe SEQ ID NO:	4019	5722	8078	7976	9251	288	870	1314		1976	4488		8684	9/06	9714	330		331	2451	2725	3073		3200	3728	4650	3463	3872	7434	7515	8732

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Single Exon Probes Expressed in Heart

											_	_										_	
Top Hit Descriptor	xf87e10.x1 NCI_CGAP_Gas4 Homo sapiens cDNA clone IMAGE:2623146 3' similar to contains MER10.t2 MER10 repetitive element;	Homo sapiens FRA3B common fragile region, cladenosine triphosphate hydrolase (FHIT) gene, exon 5	Homo sapiens hypothetical protein FLJ20585 (FLJ20585), mRNA	Homo sapiens hypothetical protein FLJ20585 (FLJ20585), mRNA	Homo sapiens FRA3B common fragile region, diadenosine triphosphate hydrolase (FHIT) gene, exon 5	Homo sapiens FRA3B common fragile region, cladenosine triphosphate hydrolase (FHIT) gene, exon 5	CANALICULAR MULTISPECIFIC ORGANIC ÁNION TRANSPORTER 1 (MULTIDRUG RESISTANCE- ASSOCIATED PROTEIN 2) (CANALICULAR MULTIDRUG RESISTANCE PROTEIN)	xb03b05.x1 NCI_CGAP_GU1 Homo sapiens cDNA clone IMAGE:2575185 3' similar to contains L1.t2 L1 repetitive element:	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG	S-ANTIGEN PROTEIN PRECURSOR	Homo sapiens LGMD2B gene	zk87a08,r1 Soares_pregnant_uterus_NbHPU Homo sapiens cDNA clone IMAGE:487858 5'	yy3c12.s1 Soares_multiple_sclerosis_2NbHMSP Homo sapiens cDNA clone IMAGE:279190 3' similar to contains L1.t3 L1 repetitive element;	wm08c03.xt NCI_CGAP_Ut4 Homo saplens cIONA clone IMAGE:2435332.3' similar to contains Alu	R.norvegicus mRNA for CPG2 protein	xp45f12.x1 NCI_CGAP_HN11 Homo saplens cDNA clone IMAGE:2743343 3' similar to contains Alu repetitive element;contains element MER9 repetitive element;	Homo sapiens a disIntegrin and metalloproteinase domain 29 (ADAM29), mRNA	hx84ft1.xf NCI_CGAP_Kid11 Home saplens cDNA clone IMAGE:31955013' similar to contains MER4.b2 MER4 repetitive element:	CIRCUMSPOROZOITE PROTEIN PRECURSOR (CS)	xp45f12.x1 NCI_CGAP_HN11 Homo sapisns cDNA clone IMAGE:2743343 3' similar to contains Alu repetitive element; contains element MER9 repetitive element;	Homo sapiens Xq pseudoautosomal region, segment 2/2	Homo saplens Xq pseudoautosomal region; segment 2/2	Homo sapiens chromosome 21 segment HS21C103
Top Hit Database Source	EST_HUMAN	Ę	N	NT	TN	FN	SWISSPROT	FST HIMAN	SWISSPROT	SWISSPROT	NT.	EST_HUMAN	EST_HUMAN	NAMI IH	LN	EST HUMAN	LN	EST HUMAN	SWISSPROT	EST HUMAN	Į.	LN	LN
Top Hit Acession No.	AW151673.1	AF020503.1	8923548 NT	8923548 NT	AF020503.1	AF020503.1	Q63120	AW073791 1	P08547	P04928	AJ007973.1	AA046502.1	N46328.1	A188924 1	X95466.1	AW265354.1	7658864 NT	BE466372.1	P02894	AW265354.1	AJ271738.1	AJ271738.1	AL163303.2
Most Similar (Top) Hit BLAST E Value	7.0E-14	6.0E-14	6.0E-14	6.0E-14	6.0E-14	6.0E-14	5.0E-14			4.0E-14	4.0E-14		4.0E-14			3.0E-14	3.0E-14	3.0E-14		3.0E-14	2.0E-14		2.0E-14
Expression Signal	3.07	10.2	1.02	1.02	2.56	2.56	3.92	90 1	5.12	1.77	6.5	0.94	-	234	2.13			1.23	1.5	7.59	3.98	3.98	6.35
ORF SEQ ID NO:		20140	24756	24757	27725	27728	20348				21608		23873		20705				24791		20154	Ц	20428
Exan SEQ ID NO:	12699	10319	14982	14982	17502	17502	10538	14860	1				14094	10780	1	l	L	14972			10331	۱ . ا	12673
Probe SEQ ID NO:	1611	363	5114	5114	7652	7652	602	4085	5397	1107	1835	3693	4194	77770	934	4841	4844	5104	5157	8563	384	384	675

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	-			Most Similar)		
Probe SEQ (D SI NO:	SEQ ID ORF ID	ORF SEQ ID NO:	Expression Signal	(Top) Hit BLAST E Value	Top Hit Acesslan No.	Top Hit Database Source	. Top Hit Descriptor
2339	12219	\dagger	1.36	2.0E-14	AW372868.1	EST_HUMAN	RC5-BT0377-091299-031-D12 BT0377 Homo sapiens cDNA
2416	12293	T	1.09	2.0E-14	7657529 NT		Homo sapiens mabdold tumor delation region protein 1 (RTDR1), mRNA
	12356	22246	1.24	2.0E-14	AL163209.2	FN	Homo sapiens chromosome 21 segment HS21,0009
1	12507		0.95	2.0E-14	P08548	SWISSPROT	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG
5605	15423	25485	2.96	2.0E-14	U01317.1	NT	Human beta globin region on chromosome 11
1	16008		2.18	2.0E-14	BE000550.1	EST_HUMAN	RC3-BN0072-240200-011-a06 BN0072 Homo sapiens cDNA
	16293	26454	19.91	2.0E-14	BE158761.1	EST_HUMAN	IL 2-HT0397-071299-024-D04 HT0397 Homo sapiens cDNA
١.		26455	19.91		BE158761.1	EST_HUMAN	IL 2-HT0397-071299-024-D04 HT0397 Homo sapiens cDNA
i		28300	4.78	2.0E-14	AW139800.1	EST HUMAN	UI-H-Bi1-adw-a-10-0-UI.s1 NCI_CGAP_Sub3 Homo sapiens cDNA clone IMAGE:2718234 3'
ı	15423	25485	1.81	2.0E-14	U01317.1	TN	Human beta globin region on chromosome 11
L	10988	20810	1.31		AL163248.2	LN	Homo saplens chromosome 21 segment HS21C046
<u>L</u>	11289	21143	7.67	1.0E-14	AL163268.2	TN	Homo sapiens chromosome 21 segment HS21,0068
	11289	21144	7.67	1.0E-14	AL163268.2	LN	Homo sapiens chromosome 21 segment HS21,0068
	-						Homo sapiens chromosome X region from filamin (FLN) gene to glucose-6-phosphate dehydrogenase
1956	11851	21738	21.54	1.0E-14	L44140.1	NT	(G6PD) gene, complete cds's
2137	12025	21921	6.17	1.0E-14	AL163303.2	NT	Homo sapiens chromosome 21 segment HS21C103
	12238	22134	6.43	1.0E-14	AF001689.1	NT	Homo sapiens ribosomal protein L23A (RPL23A) gene, complete cds
_	12841	22641	1.38	1.0E-14	P05227	SWISSPROT	HISTIDINE-RICH PROTEIN PRECURSOR (CLONE PFHRP-II)
	13055	22854	4.67		BF335227.1	EST_HUMAN	RC2-CT0432-310700-013-a09_1 CT0432 Homo saplens cDNA
	13055	22855	4.67		BF335227.1	EST_HUMAN	RC2-CT0432-310700-013-a09_1 CT0432 Homo saplens cDNA
	13723	23512	2	1.0E-14	AA682994.1	EST HUMAN	ae89c12.s1 Stratagene schizo brain S11 Homo saplens cDNA clone IMAGE:971350 3'
4374	14270	24051	1.74	1.0E-14	AW275852.1	EST HUMAN	xq38h10.x1 NCI_CGAP_Lu28 Homo sapiens cDNA clone IMAGE:2753059 3'
	_	-					Bos taurus xenobiotic/medium-chain fatty acid-CoA ligase form XL-III mRNA, nuclear mRNA encoding
6550	15466	25536	1.97	1.0E-14	AF126145.1	NI	mitochondriai protein, compiere cas
8012	19457	26045	10.41	1.0E-14	11437150 NT	. LN	Homo saplens prominin (mouse)-like 1 (PROML1), mRNA
1	19457	26046	10.41	1.0E-14	11437150 NT	NT	Homo sepiens prominin (mouse)-like 1 (PROML1), mRNA
<u> </u>	11483	21320	2.85		7427522 NT	TN	Homo sepiens protein tyrosine phosphatase, receptor type, T (PTPRT), mRNA
-	_						Homo septions transcription factor IGHM enhancer 3, JM11 protein, JM4 protein, JM5 protein, T54 protein,
200	13047		187	0 OF 15	AF106779 1	FZ	junito protein, Astalinetentatori-dependent protein, upper Lim donnan protein of and opinaproprison genes, i complete eds: and L-type calcium channel a>
6427	16288	28448	4.28	9.0E-15	P21416	SWISSPROT	GAG POLYPROTEIN [CONTAINS: CORE PROTEINS P15, P12, P30, P10]
6873	16553	26748	1.53		BE903559.1	EST_HUMAN	601677750F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3960156 5'
2780	10415		٦		8.0E-15 BE261482.1	EST_HUMAN	601148632F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:3164023 5'

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		_	_	_			-		_	_		_			_			_						_		
Top Hit Descriptor	xn77d02.x1 Soares_NFL_T_GBC_S1 Homo seplens cDNA clone IMAGE:2700483 3' similar to contains THR.t2 THR repetitive element;	Homo sapiens Xq pseudoautosomal region; segment 2/2	QV1-LT0036-150200-070-c10 LT0036 Homo sapiens cDNA	nab81c12.x1 Soares_NSF_F8_9W_OT_PA_PIS1 Homo sapiens cDNA clone IMAGE: 3'	Homo sapiens chromosome 21 segment HS21C008	Human hereditary haemochromatosis region, histone 2A-like protain gene, hereditary haemochromatosis (HLA+I) gene, RoRet gene, and sodium phosphata transporter (NPT3) gene, complete cds	UI-H-BW0-qjb-g-10-0-UI.s1 NCI_CGAP_Sub6 Homo sapiens cDNA clone IMAGE:2731219 3'	AV730056 HTF Homo sapiens cDNA clone HT FAVE06 5	Homo saplens chromosome 21 segment HS21C103	DKFZp761C0810_r1 761 (synonym: hamy2) Hömo sapiens cDNA clone DKFZp761C0810 51	Homo sapiens mRNA for transcription factor	Homo sapiens mRNA for transcription factor	LY1142F Human fetal heart, Lambda ZAP Express Homo sepiens cDNA clone LY1142 5' similar to ANF(CARDIODILATIN)	NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 5	7P01F03 Chromosome 7 Placental cDNA Library Homo saplens cDNA clane 7P01F03	7P01F03 Chromosome 7 Placental cDNA Librery Homo saplens cDNA clone 7P01F03	Mus musculus ultra high sulfur keratin gene, complete cds	Mus musculus ultra high sulfur keratin gene, complete cds	oc38a07.s1 NCI_CGAP_GCB1 Homo sapiens oDNA clone IMAGE:1351764.3' similar to contains MER19.t1 MER19 repetitive element;	Homo saplens DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes,	complete cds)	Homo sapiens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively spliced.	Homo capiene calcium channel alpha TE subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively	spilose	Homo sapiens calcium charnel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively spliced	Homo sapiens hypothetical protein FLJ20212 (i*LJ20212), mRNA
Top Hit Database Source	EST_HUMAN	Z L	EST_HUMAN	EST_HUMAN	Z	L	EST HUMAN	EST HUMAN	N	EST_HUMAN	Z	Z	EST HUMAN	SWISSPROT	EST_HUMAN	EST_HUMAN	TN	LN	EST HUMAN		L	TN	!	-Z	TN	TN
Top Hit Acession No.	AW241958.1	AJ271736.1	AW 836843.1	BF432200.1	AL163208.2	U91328.1	AW296817.1	AV730056.1	AL163303.2	AL118596.1	AJ130894.1	AJ130894.1	N89452.1	P92485	AA078097.1	AA078097.1	M27685.1	M27685.1	AA807128.1		AB026898.1	AF223391.1		AF223391.1	AF223391.1	8923201 NT
Most Similar (Top) Hit BLAST E Value	7.0E-15	6.0E-15	6.0E-15	6.0E-15	5.0E-15	5.0E-15	6.0E-15	5.0E-15	4.0E-15	4.0E-15	4.0E-15					3.0E-15		3.0E-16	3.0E-15		3.0E-15	2.0E-15	_	2.0E-15	2.0E-15	2.0E-15
Expression Signal	2.83	6.12	1.86	1.67	5.79	1.38	F	2.22	2.6	0.78	238	2.38	5.93	1.41	0.88	0.88	2.86	2.86	1.87		2.71	3.29		3.23	3.23	1.14
ORF SEQ ID NO:		20748			20177	22490			19779	23720	26623				24589		26335	26336			28311	20033	_	20141	20142	
Exen SEQ ID NO:	17813	10901		19415	10350	12595	13340	17954	8866	13942	16438	16438	14023	14724	14832	14832	16177	16177	17559		18061	10216		10320	10320	11417
Probe SEQ ID NO:	7963	878	8622	8948	404	2733	3423	8063	421	4039	8414	8414	4123	4843	4955	4955	6314	6314	7709		8173	250		384	364	1512

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Table 4
Single Exon Probes Expressed in Heart

_					_			_		_	_	_	_		_			_	_	-	_			_	_		_	
	Top Hit Descriptor	Homo sapiens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively spliced	Homo saplens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively spliced	wf07f06.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2349923 3' similar to TR:Q61043 Q61043 NINEIN :	REPETITIVE PROLINE-RICH CELL WALL PROTEIN 2 PRECURSOR	REPETITIVE PROLINE-RICH CELL WALL PROTEIN 2 PRECURSOR	Homo sapiens ASCL3 gene, CEGP1 gene, C11orf14 gene, C11orf15 gene, C11orf16 gene and C11orf17 gene	2/77e03.s1 Scares_fetal_fiver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:460924 3'	za78410.r1 Scares_fetal_jung_NbHL19W Homo saplens cDNA clone IMAGE:298875 5' similar to WP:E44F4 8 CE02227 TRANSPOSASE:	Human DNA, SINE repetitive element	CM0-HT0244-201099-078-a12 HT0244 Homo sapiens cDNA	CM0-HT0244-201099-078-e12 HT0244 Homo saplens cDNA	Homo capiens Xq pseudoautosomal region; segment 1/2	Homo sapiens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively	Springer Londo concieno exterior absorbed plabade submit (CACAIAE) was extens 7.40 and nortical ads. allegantingly	Thurs definite countril triannel apria in countril (Society Society) and partial countries and partial countries and partial countries and partial countries and countries are considered and countries and countrie	bz6h05x1 NCI_CGAP_Lu24 Homo saplens cDNA clone IMAGE:2270745 3' similar to TR:Q13539 Q13539	hk40e02.vf NCI CGAP Ov34 Home sapiens ci0NA clone IMAGE:2999162 5'	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG	RC3-HT0649-100500-022-b05 HT0849 Homo sapiens cDNA	wr86e04.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2494590 3'	ye40e10.51 Soares fetal liver spleen 1NFLS Horno sapiens cDNA clone IMAGE:120234 3' similar to contains	MER6 repetitive element;	QV3-BT0569-270100-074-g05 BT0569 Homo saplens cDNA	Homo saplens chromosome 21 segment HS21C080	qf68h08.xt Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1755227 3'	qf88h06.x1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1755227 3'	Homo sapiens spermidine synthase (SRM) mRNA
	Top Hit Database Source	Į,	Į.	EST_HUMAN	SWISSPROT	SWISSPROT	ΙΝ	EST_HUMAN	FST HUMAN	FN	EST_HUMAN	EST_HUMAN	N	ţ	2	뒫	TO L	EST HUMAN	SWISSPROT	EST_HUMAN	EST_HUMAN		EST_HUMAN	EST_HUMAN	N	EST_HUMAN	EST HUMAN	TN
	Top Hit Acession No.	AF223391.1	AF223391.1	AI806335.1	P13993	P13993	AJ400877.1	AA704185.1	W05084 1	D14547.1	AW379465.1	AW379465.1	AJ271735.1	A Engage 4	AT 260391.1	AF223391.1	4 4000001 A	BE043584.1	P08547	BE182696.1	Al984928.1		T95763.1	BE074217.1	AL163280.2	AI200976.1	AI200976.1	4507208 NT
	Most Similar (Top) Hit BLAST E Value	2.0E-15	2.0E-15	2.0E-16	2.0E-15	2.0E-15		2.0E-15				2.0E-15	2.0E-15	2 DO C	Z.UE-13	2.0E-15	4 00 45				1.0E-15		1.0E-15	1.0E-16			1.0E-15	1.0E-15
	Expression Signal	1.04	1.04	2.07	1.33	1.33	1.71	2.2	513				3.01		77.7	2.22		80					1.83	1.98	1.28		4.67	1.44
	ORF SEQ ID NO:	23188	23187		24741			26331		L	27484				23100	23187		22706		L	24731		25864					27293
	Exon SEQ ID NO:	13381	13381	14415	14866	14966	16089	16173	18237	J.	17271	17277	18096	10001		13381	7.0000	12908)	ı	l		15750	16068	16670		16783	1
	Probe SEQ ID NO:	3465	3465	4522	2087	2097	6223	6310	6376	7163	7410	7410	8212	0420	SAIA	8189	1	2978	3103	4262	5086		5844	6182	6791	9069	9069	7227

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Single Exon Probes Expressed in Heart

Most Similar Expression (Top) Hit Signal BLAST E Value	8331 6.81 1.0E-15 AF044083.1 NT Homo sapiens major histocompatibility locus class III region	tr31c05.x1 NCI_CGAP_Ov23 Homo sapiens cDNA clone IMAGE:2219912.3' similar to contains Alu repetitive from the first from the f	1.03 9.0E-16 4503168	2.6 9.0E-16 F08688.1	8388 1.6 7.0E-16 O88807 SWISSPROT (PEPTIDYLARGININE DEIMINASE TYPE ALPHA)	5,1	6.8 7.0E-16 T94149.1 [EST_HUMAN ye28c12.r1 Stratagene lung (#937210) Homo sapiens cDNA clone IMAGE:119062 5'	8.32 6.0E-16 AW972611.1 EST_HUMAN EST384702 MAGE resequences, MAGL Homo sapiens cDNA	1.09 5.0E-16 AJ251154.1 NT Mus musculus olfactory receptor cluster, OR37A, OR37E, OR37E genes and OR37D pseudogene	0180c04.81 Scares_total_fetus_Nb2HF8_9w Home sapiens cDNA clone IMAGE:1623078 3' similar to	4 60 6 46 At 489046 0	3.33 6.0E-16 BF217368.1 EST HUMAN	8.34 5.0E-16 11418127 NT	1.27 4.0E-16/AB001523.1 NT Homo sepiens gene for TMEM1 and PWP2.complete and partial cds	1.32 4.0E-16 AW 797168.1 EST_HUMAN	1.32 4.0E-16 AW 797168.1 EST_HUMAN	3.85 4.0E-16 Q16853 SWISSPROT	3.55 4.0E-16 BE083875.1 [EST_HUMAN	3.55 4.0E-16 BE083875.1 EST_HUMAN	1 4.0E-16 P08548 SWISSPROT	35.8 4.0E-16 AL163284.2 NT Homo sepiens chromosome 21 segment HS21,0084	1.22 4.0E-16 11423191 NT	1.74 4.0E-16[AV730030.1 EST_HUMAN	P08548 SWISSPROT	5,94 4.0E-16 C05947.1 EST_HUMAN C05947 Human pancreatic islet Homo capiens cDNA clone hbc5355	2.04 4.0E-16 6912459 NT	1.59 3.0E-16 AW022862.1 EST_HUMAN	1.59 3.0E-16 AW022862.1 EST_HUMAN	1,47 3.0E-16/AL049445.1 EST_HUMAN DKFZp434P037_r1 434 (synonym: htes3) Homio sapiens cDNA clone DKFZp434P037 5'
	6.81	3.71	1.03	26	1.5	5.	6.8	8.32	1.09	7.07	6/:	3.33	8.34	1.27	1.32	1.32	3.85	3.55	3.55	1	33.8	1.22	1.74	1.64	5.94	2.04	1.59	1.59	1.47
o B O	18079 28331	19492 25131	14298 24082	18238 28486	16206 26368	16206 26369	19585	11983	11382 21246	1351	1	18623 28914	19381	12079	12209 22107				13952 23729	14962 24737	16396 26575	17226 27425	18421 28691	18906	18957	18964 25319	10102 19924	10102 19925	10402
Probe Exan SEQ ID SEQ ID NO: NO:	8183 18	9867 19	4404 14	8361 18	6343 16		9816 19		1477 11	7,790	L	8809 18	L	2192 12							. '				9244 18	9255 18	128 10	128 10	458 10

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	Top Hit Descriptor	Homo sapiens pitultary tumor transforming gene protein (PTTG) gene, complete cds	QV0-0T0032-080300-155-d01	Homo sapiens chromosome 21 segment HS21C080	MR0-HT0559-060300-003-e04 HT0559 Homo sippiens cDNA	AV730759 HTF Homo saplens cDNA clane HTFAQB07 5'	Mus musculus apolipoprotein B editing complex 2 (Apobec2), mKNA	Homo sapiens putative MTAP (MTAP) mRNA, partial cds, alternatively spliced	Mus musculus WNT-2 gene, partial cds; putativi ankyrin-related protein and cystic fibrosis transmembrane	conductance regulator (CFTR) genes, section 1 of 2 of the complete cds; and unknown gene	RC1-HN0003-220300-021-b04 HN0003 Homo sapiens cDNA	hi81404.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2978695 3 similar to contains L1.tZ L1 repetitive element ;	yc05h08.r1 Stratagene lung (#937210) Homo sepiens cDNA clone IMAGE:79839 5'	yd26b04.r1 Soares fetal liver spleen 1NFLS Homo saplens cDNA clone IMAGE:109327 5'	Homo sepiens chromosome 21 segment HS21C047	ov45e04.x1 Soares_tests_NHT Homo sapiens cDNA clone IMAGE:1640286 3' similar to TR:Q16530	Q16530 PMS3 MRNA ;contains MER10.t2 MER10 repetitive element;	xd89c09.x1 Soares_NFL_T_GBC_S1 Homo sapiens cONA clone IMAGE:2004/84 3	MAS-RELATED G PROTEIN-COUPLED RECEPTOR MRG	hw05b04.xf NCI_CGAP_Lu24.Homo sapiens cDNA clane IMAGE:31318989.3	hw05b04 x1 NCI_CGAP_Lu24 Homo sapiens cUNA clone IMACE:31818989 3	Homo sapiens DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, ORC1L3, ORC1L4 genes,	complete cds)	Hamo saplens SEC14 (S. cerevisiae)-like 2 (SEC14L2), minna	AV720204 GLC Homo sapiens cDNA clone GLCDIF08 5	qt63a06.x1 NCI_CGAP_Eso2 Homo sapiens ct0NA clone IMAGE:1959922 3' similar to contains Alu	Tepenuve exertion, Les Homes enjoye cityld clane (MAGE-1959223' similar to contains Alu	groadoux i Not_confEsta i unio saprena con o ciclo monte.	zg81d04.s1 Soares_fetal_heart_NbHH19W Homo sapiens cDNA clone IMAGE:399751 3'	ZONADHESIN PRECURSOR	ZONADHESIN PRECURSOR
TopHit	Database Source	NT	EST_HUMAN	INT	EST_HUMAN	EST_HUMAN	닐	NT		NT	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	Ν		EST_HUMAN	EST_HUMAN	SWISSPROT	EST_HUMAN	EST_HUMAN		۲	INT	EST HUMAN		ESI HUMAN	EST HUMAN	EST_HUMAN	SWISSPROT	SWISSPROT
	l op Hit Acession No.	AF200719.1	AW880701.1	AL 163280.2	BE172081.1	AV730759.1	TN 7605379	AF216850.1		AF229843.1	AW983880.1	AW682772.1	T64110.1	T81043.1	AL163247.2		A1073546.1	AW119123.1	P35410	BE326522.1	BE326522.1		AB02689		AV720204.1		AI270080.1	AI270080.1			Q28983
N	(Top) Hit BLAST E Value	9.0E-17	8.0E-17		8.0E-17	8.0E-17	7.0E-17	_		7.0E-17		R 0E-17	_				4.0E-17	3.0E-17	3.0E-17	3.0E-17	3.0E-17		3.0E-17	3.0E-17	3.0E-17		2.0E-17	2.0E-17		2.0E-17	2.0E-17
	Expression Signal	2.18	1.7	0.87	3.56	1.36	3.18	3.05		6.83	4.78	18.4	234	207	2.12		1.98	1.35	1.31	1.14	1.14		4.72	3.16	15.82		2.81	- 8	1.27	2.6	2.6
	ORF SEQ ID NO:				26402					26052		08830				1		21836		23283			27680				20127	20127		22170	
, A	SEQ ID	17720	10919	13729	19444	16174	11347			16921		i .	L	ľ	1	ì	18914	11941	L	13504	l_	L_	17445	ł_	L	<u> </u>	10309	10309	L	Ł	Ш
Grapha	SEQ ID	7870	1001	3817	5427	6311	1442	5262		6017	188	200	2001	2778	8783		9171	2051	3157	3590	3590		7594	9134	88		350	35,	972	2397	2397

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Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Vætue	Top Hit Acession No.	Top Hit Database Source	Tóp Hit Descriptor
2899	12826	22621	5.62	2.0E-17	P12036	SWISSPROT	NEUROFILAMENT TRIPLET H PROTEIN (20) KDA NEUROFILAMENT PROTEIN) (NEUROFILAMENT HEAVY POLYPEPTIDE) (NF-H)
5295	15216		1.95	2.0E-17	M27685.1	F	Mus musculus ultra high suifur keratin gene, complete cds
5235	15216	25018	1.95	2.0E-17	M27685.1	ΙN	Mus musculus ultra high sulfur karatin gene, complete cds
5790	15696		2.07	2.0E-17	AF055066.1	FZ	Homo sapiens MHC class 1 region
6711		26779	1.44		Q95158	SWISSPROT	OLFACTORY RECEPTOR-LIKE PROTEIN OLF3
6925	16803		1.38	20E-17	AA300640.1	EST_HUMAN	EST13504 Testis tumor Homo saplens cDNA 5' end similar to similar to glycogenin
7699	17549	27772	2.72	20E-17	AL163247.2	LN	Homo saplens chromosome 21 segment HS210047
7699	17549	27773	2.72	2.0E-17	AL163247.2	LN	Homo sapiens chromosome 21 segment HS210047
							Human CYP19 gene for aromatase cytochrome P-450, promoter region (containing two cis-ecting
7887			5.3	2.0E-17		NT	transcriptional regulatory elements)
733		20499	3.37	1.0E-17	P08183	SWISSPROT	MULTIDRUG RESISTANCE PROTEIN 1 (P-GLYCOPROTEIN 1)
1729			2.45	1.0E-17	AL163207.2	NT	Homo saplens chromosome 21 segment HS210007
2069			1.68	1.0E-17	P02461	SWISSPROT	COLLAGEN ALPHA 1(III) CHAIN PRECURSOR
2287	12170		1.48	1.0E-17	U79410.1	TN	Homo saplens thrombospandin 2 (THBS2) gene, promoter region and exons 1A and 1B
						•	Homo sapiens mannosidase, beta A, lysosomal (MANBA) gene, and ubiquitin-conjugating enzyme E2D 3
3519	١		1.01	1.0E-17	9.1	NT	(UBE2D3) genes, complete cds
4045	13947		7.17	1.0E-17	R09942.1	EST_HUMAN	y/30e07.r1 Soares fetal liver spleen 1NFLS Hono sapiens cDNA clone IMAGE:128388 5'
							he38e05.x1 NCI_CGAP_CML1 Homo sapiens cDNA clone IMAGE:2921312 3' similar to contains Alu
5885			4.7		AW468468.1	EST_HUMAN	repetitive element; contains LTR8.t1 LTR8 repetitive element;
9009		26037	1.44			EST HUMAN	qe65b05.x1 Soares_fetal_lung_NbHL19W Homo sapiens cDNA clone IMAGE:17438253'
9009			1.44		AI185642.1	EST_HUMAN	qe63b05.x1 Soares_fetal_lung_NbHL19W Homo sapiens cDNA clone IMAGE:1743825 3'
6216			1.32		Q16831	SWISSPROT	URIDINE PHOSPHORYLASE (UDRPASE)
8719			2.01	1.0E-17	Q28824	SWISSPROT	MYOSIN LIGHT CHAIN KINASE, SMOOTH MUSCLE (MLCK) [CONTAINS: TELOKIN]
2422		22196	96.0	9.0E-18	AA174078.1	EST_HUMAN	zp18g12.s1 Stratagene fetal retina 937202 Homo saplens cDNA clone IMAGE:609862 3'
7468			3.26	9.0E-18	AI472167.1	EST_HUMAN	tased03.x1 Soares_NSF_F8_9W_OT_PA_P_S1 Hamo sapiens cDNA clone IMAGE:2148389 3'
3718	13630	23415	1.75	8.0E-18	4758977 NT	LN	Homo sapiens protein tyrosine phosphatase, non-receptor type substrate 1 (PTPNS1) mRNA
							xx10b04,x1 NCI_CGAP_Pan1 Homo saplens cDNA clone IMAGE:2837071 3' sImilar to gb:L20868 60S
348	10305	20121	8.39	7.0E-18	AW316976.1	EST_HUMAN	RIBOSOMAL PROTEIN L4 (HUMAN);
346	10305	20122	8.39	7.0E-18	AW316976.1	EST_HUMAN	xx10b04.x1 NCI_CGAP_Pen1 Homo sapiens cDNA clone IMAGE:2837071 3' similar to gb:L20868 60S RIBOSOMAL PROTEIN L4 (HUMAN);
	!				,		ya49c07.s1 Soares Infant brain 1NIB Homo sapiens cDNA clone IMAGE:53285 3' similar to contains L1
5136	15003	24774	0.85	7.0E-18	R16220.1	EST_HUMAN	repetitive element ;

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Single Exon Probes Expressed in Heart

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acesslan No.	Top Hit Database Source	Tap Hit Descriptor
8638	10305	20121	5.73	7.0E-18	AW316976.1	EST_HUMAN	xx10b04.x1 NCI_CGAP_Pan1 Homo sapiens cCiNA clone IMAGE:2837071 3' sImilar to gb:L20868 60S RIBOSOMAL PROTEIN L4 (HUMAN);
8838	L		6.73	7.0E-18	AW316976.1	EST_HUMAN	xx10b04.x1 NCI_CGAP_pan1 Homo sapiens cDNA clone IMAGE:2837071 3' similar to gb:L20868 60S RIBOSOMAL PROTEIN L4 (HUMAN);
3255	L	22976		6.0E-18	X71791.2	Z	Rattus norvegicus partial Gdn/Pn-1 gene for glia-derived nexin/protease nexin I, enhancer region
4641	Ĺ		3.37	6.0E-18	P52181	SWISSPROT	PROTEIN-GLUTAMINE GAMMA-GLUTAMYLTRANSFERASE (TISSUE TRANSGLUTAMINASE) (TGASE C) (TGC)
6789	1		2.69			Į į	Homo sapiens similar to high-mobility group (norihistone chromosomal) protein 4 (H. sapiens) (LOC63446), mRNA
8476	ſ	28614			AL1632	NT TA	Homo saplens chromosome 21 segment HS21C046
8642	1				X87344.1	Į.	H.sapiens DMA, DMB, HLA-Z1, IPP2, LMP2, TAP1, LMP7, TAP2, DOB, DQB2 and RING8, 9, 13 and 14 genes
9394			3.66	81-30E	U87929.1	N.	Human aconitate hydratase (ACO2) gene, exon 4
1130	l				AI280214.1	EST HUMAN	qm65g11.x1 Soares_placenta_8to9weeks_2NbHP8to9W Homo sapiens cDNA clone IMAGE:1893668 3' similar to contains Alu repetitive element;
4217	1					NT	Mus musculus gasdermin (Gsdm), mRNA
5058				ļ	D61517	EST HUMAN	HUM411F05B Ciontech human fetal brain pdyA+ mRNA (#6535) Homo sapiens cDNA clone GEN-411F05 5
5224					_	N	Human endogenous retrovirus HERV-P-T47D
7061	L					EST_HUMAN	MR0-HT0161-221099-002-c06 HT0161 Hamo caplens cDNA
8346	<u> </u>		4.33	5.0E-18	10242378 NT	TN	Homo sapiens lymphocyte activation-associated protein (LOC51088), mRNA
8346	L			5.0E-18	10242378 NT	Ā	Homo sapiens lymphocyte activation-associated protein (LOC\$1088), mRNA
9512	L				AW867182.1	EST_HUMAN	MR1-SN0035-060400-001-g11 SN0035 Homo saplens cDNA
9837	ł		13	5.0E-18	AV650547.1	EST_HUMAN	AV650547 GLC Homo sapiens cDNA clone GLCGA02 3'
	10096	18915	1.36	4.0E~18	BE044076.1	EST HUMAN	ho36h04.x1 NCI_CGAP_Ut1 Homo sapiens cDNA clone IMAGE:3039511 3' sImilar to contains MER29.b3 MER29 repetitive element ;
	i				_	HOT LON	ho38h04.x1 NC_CGAP_Ut1 Homo sapiens cDNA clone IMAGE:3039611 3' similar to contains MER29.b3 MER29 canadition element
119		18916				NEW TOTAL	Military operator operators of the Home consister of TAIA close IMAACE 2302005 3'
1846	11742		1.08	4.0E-18	AI738592.1	ESI HUMAN	WISSINGS I NOT CASH. COLD Train Saprens Civil Colors (NOT CASH)
. 2164	12042	21840	0.98	4.0E-18	Q06430	SWISSPROT	N-ACE I'LLACI IOSAMINIDE BE IA-1, S-Y-ACE I TL'SLOCOSAMINIT I RANSFERASE (N- ACETYLGLUCOSAMINYL TRANSFERASE) (I;BRANCHING ENZYME) (IGNT)
2154	12042	21841	0.98	4.0E-18		SWISSPROT	N-ACETYLLACTOSAMINIDE BETA-1,6-N-ACETYLCLUCOSAMINYLTRANSFERASE (N- ACETYLCLUCOSAMINYLTRANSFERASE) (I;BRANCHING ENZYME) (IGNT)
6233	15214	}	2.55	4.0E-18	AI017565.1	EST_HUMAN	ou23e06.x1 Soares_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:1627138 3'

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Single Exon Probes Expressed in Heart

Top Hit Descriptar	ou23e06x1 Scares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:162/138 3	EST 69633 Printery gland, subtracted (protecting/own namore) in the containing of anily repeat EST containing of family repeat	ob23h11:s1 NCI_CGAP_Nos normo septens curva cignis in curva cignis in curva cignis in curva cignis cignis curva cignis cigni	CM0-BT0690-210300-298-g07 B1099 Homo capiens culting	Homo sapiens chromosome 21 segment noc rough	PM0-BN0081-100300-001-006 BN0061 notice spaces out A	113.10.12. yi Morton retal Cocilies notific september China.	QV1-L 10030-150Z0U-0/0-e0/ L10050 monitors serving	601114352F1 NIH MIGC TO FIGURE SEDIM CONTROLL SOCIETY SIGNIFIER TO TR 014577	ak33a07.51 Soares, tests, Nr1 India septetts cure information of the cure information of the cure re-14406 FROM 7031, COMPLETE SEQUENCE.	Human DNA, SINE repetitive element	Human DNA, SINE repetitive element	602021164F1 NCI_CGAP_Bm67 Homo sapiens cDNA clone IMAGE:4130610 3	hig4g01.x1 Soares_NFL_T_GBC_S1 Hamb sapiens cDNA clone invAGE_2313304.3 Strings to Strings to MER19.12 MER19.12 MER19 repetitive element;	ARZA10 VI NCI CGAP Gas4 Homo sapiens cDNA clone IMAGE: 2623146 3' similar to contains MER10.12	MER10 repetitive element;	x67e10.x1 NCI_CGAP_Gas4 Homo sapiens cDNA clone IMAGE:2023140 3 Similar to contains with 19.22	MERTO repeature cientral .	has3a06.x1 NCI_CGAP_NIG12 Homo seprens color non-color color color nancolor color color nancolor color nancolor color nancolor color nanco	xg47e09.x1 NCI_CGAP_Utf Homo sapiens cDNA clone IMAGE:2630728 3' similar to contains MER8.b2	MER8 repetitive element;	601114352F1 NIH_MGC_16 Homo saplens cDlyA clone IMAGE:3355044 5	ye43g05.r1 Soares fetal liver spieen 1NFLS Homo sapiens cDNA clone IMAGE:120536 5' similar to contains	L1 repetitive element;	AV653405 GLC Homo sapiens cDNA clone GLCDKE11 3	Homo sapiens mRNA for Na,K-ATPase alpha-subunit, complete cds	Homo sapiens mRNA for Na,K-ATPase alpha-subunit, complete cds	Homo sapiens chromosome 21 segment HSZI (3080
Top Hit Database Source	EST_HUMAN	EST_HUMAN	HUMAN	EST_HUMAN		T	Т	7	EST_HUMAN	EST_HUMAN	TN	NT	EST_HUMAN	NAM HI MAN		EST_HUMAN		EST_HUMAN	EST HUMAN	II.	EST_HUMAN	EST HUMAN		EST_HUMAN	EST HUMAN	ᅜ	NT	NT
Top Hit Acession No.	AI017565.1	AA371807.1			AL163247.2				BE256097.1	AA868610.1	D14547.1	D14547.1	BF347229.1	A18/06/50 4	-	AW151673.1	_	AW151673.1	AW470791 1		AW151299.1	BE256097.1		T95408.1	AV653405.1	D00099.1	D00099.1	AL163280.2
Most Similar (Top) Hit BLAST E Value	4.0E-18 A	4.0E-18	3.0E-18	3.0E-18	3.0E-18	3.0E-18	3.0E-18	2.0E-18	2.0E-18	2.0E-18	2.0E-18	2.0E-18	2.0E-18	04	Z.UE-10	2.0E-18		2.0E-18	9.E-18	L		2.0E-18		1.0E-18	L	L	L	Ш
Expression Signal	2.55	7.12	2.38	2.41	1.19	5.2	4.92	2.83	47.22	3.2	3.04	3.04	1.67		3.55	1.83		1.63	68.9	2000	4.44			1.02				
ORF SEQ ID NO:	25015	28489	20609	L				20034			25149				25834	27861	L	27862			29085				24985			Ш
Ewn SEQ ID NO:	15214	18247	10758	10838	13778	16029	19210	10217	11049	15242	1	1	1	1	15720	17629	1	17629	l	1921/	18774	1		14215	1	1		1 1
Probe SEQ ID NO:	5293	8370	831	914	3867	6084	9842	251	1135	5322	5380	5380	5585		5814	7770		7779	3	834C	8908	9000	222	4318	5286	5440	5418	5883

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Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
7696	17546	27769	4.4	1.0E-18	U91328.1	FN	Human hereditary haemochronatosis region, histone 2A-ilke protein gene, hereditary haemochromatosis (HLA-H) gene, RoRet gene, and sodium phosphate transporter (NPT3) gene, complete cds
8277	18980	25324	2.53	1.0E-18	AF003529.1	TN	Homo sapiens glypican 3 (GPC3) gene, partial cds and flanking repeat regions
533	10475	20289	4.28	9.0E-19	ÅA281961.1	EST_HUMAN	z11d06.r1 NCI_CGAP_GCB1 Homo sapiens cINA clone IMAGE:712811 5' similar to contains MER19.t2 MER19 repetitive element;
534	. 10475	20289	3.19	9.0E-19	AA281981.1	EST HUMAN	z11d06.r1 NCI_CGAP_GCB1 Homo sapiens cIDNA clone IMAGE:712811 6' similar to conteins MER19.t2 MER19 repetitive element;
9284	_		6.21	9.0E-19	F08688.1	EST_HUMAN	HSC23F051 normalized infant brain cDNA Homb saplens cDNA clone c-23f05
7042	16919	27110	2.46	9.0E-19	AL163203.2	TN	Homo sapiens chromosome 21 segment HS210003
7042		27111	2.48		AL163203.2	NT	Hamo saplens chromosome 21 segment HS21C003
8470	18343	28608	3.88	9.0E-19	AB032969.1	TN	Homo sapiens mRNA for KIAA1143 protein, partial cds
9042	10475	20289	8.69	9.0E-19	AA281961.1	EST HUMAN	z11d06.r1 NCI_CGAP_GCB1 Homo eapiens cDNA clone IMAGE:712811 5' similar to contains MER19.t2 MER19 repetitive element ;
1032	10950		1.54	8.0E-19	AW974902.1	EST_HUMAN	EST387007 MAGE resequences, MAGN Homo, sapiens cDNA
2198	12085	21987	1.58	7.0E-19	4758139 NT	TN	Homo sapiens DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 6 (RNA helicase, 54kD) (DDX6) mRNA
5884	15791	25913	2.15	7.0E-19	AF092090.1	TN	Rattus norvegicus cp151 mRNA, partial cds
9179	19742		2.95	7.0E-19	AA705684.1	EST_HUMAN	260b01.s1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:435145 3'
3713			1.02	6.0E-19	AW852830.1	EST_HUMAN	PM0-CT0248-131099-001-g01 CT0248 Homo capiens cDNA
4360	14258	24041	1.38	6.0E-19	P34986	SWISSPROT	OLFACTORY RECEPTOR 6 (M50)
4360	14256	24042	1.36			SWISSPROT	OLFACTORY RECEPTOR 6 (M50)
4694	14580		1.18	6.0E-19	AJ271735.1	LNT	Homo sapiens Xq pseudoautosomal region; seginent 1/2
4952	14829	24595	1:09	6.0E-19	AL120817.1	EST_HUMAN	DKFZp782F192_r1 782 (synonym: hmel2) Homb sapiens cDNA clone DKFZp762F192 5'
5571	15486	25562	5.24	5.0E-19	000193	SWISSPROT	ZONA PELLUCIDA SPERM-BINDING PROTEIN B PRECURSOR (ZONA PELLUCIDA GLYCOPROTEIN [ZP-X] (RC55)
8824			7.19			EST HUMAN	x87b02xt Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:28641713' similar to contains element MSR1 repetitive element;
							Human germline T-cell receptor beta chain TORBV13S1, TORBV6S8A2T, TORBV5S6A3N2T, TORBV13S6A2T, TORBV6S9P, TORBV5S3A2T, TORBV13S8P, TORBV6S9P, TORBV13S4, TORBV5S2A1N1T, TORBV5SAA2T, TORBV6SSAA1
9849	18495		1.52	5.0E-19	U66060.1	NT	TCRBV23S1A2T, TCRBV12>
542	l		1.45			TN	Homo sapiens mRNA, chromosome 1 specific transcript KIAA0501
2849	12516	22406	1.39	4.0E-19	BF697362.1	EST_HUMAN	602130910F1 NIH_MGC_56 Home saplens cDINA clone IMAGE:4287674 5'

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				Most Similar		789 Hit	
Probe SEQ ID NO:	SEQ ID	ORF SEQ ID NO:	Expression Signal	(Top) Hit BLAST E Value	Top Hit Acesslon No.	_	Top Hit Descriptor
3778	13690	23476	1.74	3.0E-19	128997		BETA-2 ADRENERGIC RECEPTOR
3778	ì		1.74		128997		BETA-2 ADVENERGIC RECEPTION
4345	1.		1.18	3.0E-19	1V708136.1	EST_HUMAN	AV708136 ADC Homo saptens CLINA Claris AUCANIVATI O
	L						Homo sapiens similar to aldo-keto reductase tarrilly 1, member B i 1 (aldosa reductase-line) (* 1. Septembrilly 1. Member B i 1 (aldosa reductase-line) (* 1. Septembrilly 1. OC83222), mRNA
6370	16232			3.05-19	11432214		M. Consider and M. For TDOR33 modeln
7443	16456	26645			X89685.1	2	M. musculus III Alva III Oraci process
9416	19068		7.38		AF165520.1	2	Homo septers protocing in configurations and analysis of configurations and configuration
2513	12387				AL163201.2	П	Homo sapiens circumosonie za seguinari nostrador de lona IMAGE 360880 6.
6849	L	26923	8.96	2.0E-19	AA012854.1	╕	Z834GUSLT Socials require No. September Septem
473	10417	L	1.8	1.0E-19	BE408611.1	EST_HUMAN	60130412511 NIH MCC_ZI Indian september Color into Contains
	į .				100705 4	NAMILL FAST	yo79g07.r1 Soares adult brain N2b4Hb557 Homo sapiens con A cigne invoce. For 50 continue a commercial management.
2118	1	21906			T30/83.1	F14	Hirman gene for Ah-receptor, exon 7-9
2685			2.88	1.0E-19	D38044.	Į.	Thems sales make in brossine phosphatase, non-receptor type substrate 1 (PTPNS1) mRNA
2817	12746	3	5.03		4758977 NI	Z	Truth September 19 and
٤	72007	22026	1.01	1 0È-19	AA834967.1	EST HUMAN	graph.zsi Soares Jesus Jin in nomo sapidno son visita de la companya de la compan
3 3						LN	Oryctolagus cuniculus sodium/dicarboxy/ate cotransporter mRNA, partial cds
	1				MARARY 1	LΝ	Rabbit phosphorylase kinase beta subunit mRNA, complete cds
88	16797	7,0550	2.7		1.100011		w72h02.1 Soares fetal liver spleen 1NFLS Horno sapiens cDNA clone IMAGE:123243 5' similar to contains
7076	16953		2.83		T99920.1	EST_HUMAN	OFR repetitive element;
7853	1	3 27948	3 25.21	1.0E-19	AW812259.1	EST_HUMAN	RC0-S101/4-191099-031-003-0101/4-101/10 capteris contact
7857	L	L	1.79	1.0E-19	N44631.1	EST HUMAN	ly31e09.r.1 Soares melanocyte Zivonivi nollilo sapratis duriva cica melanocyte Zivonivi nollilo sapratis duriva cica menanocyte Zivonivi nollilo sapratis duriva cica menanocyte zivonivi nollilo sapratis di cica menanocyte zivonivi nollilo sapratis di cica menanocyte zivonivi nollilo sapratis di cica menanocyte di cica menanocyte di cica menanocyte di cica menanocyte di cica menanocyte di cica menanocyte di cica menanocyte di
6005	L	26035	5 2.22			NT	Mus musculus keretin-associated protein 9-1 (ni labor-1), minno
6005	L					NT	Mus musculus keratin-associated protein 9-1 (r) taber 1), mistro
6439	L		1.31	1 8.0E-20		EST HUMAN	I GORGOUGIXT SORRES INTL. I GOO OF HOME SOURCE CONTRIBUTE OF THE INVAINT HOME INVAINTS OF THE
6439		0 26463	1.31		AI221371.1	EST HUMAN	Igg86709.x1 Soares_NrL_I_GEC_31 Anome Lamp confers CONA
3238	L	L	1 0.88		BF326455.1	EST_HUMAN	PM4-AN0096-050900-003-804 AN0090 TOTO SEQUENCE CONT.
8178	1.			3 7.0E-20	AL138120.1	EST_HUMAN	DKFZp5470092_r1 547 (synonym: mpn) homo septens curve come to the control of the
							nideco4.s1 NCI_CGAP_Pr4 Homo sapiens cDNA clone IMAGE:1043/16 similar to contains interested
6947	16825	5 27018	10.99	9 7.0E-20	0 AA557657.1	EST HUMAN	MER29 repetitive element;
	1				A A C C C C C C C C C C C C C C C C C C	EDT E	MER29 repetitive element;
6947	1	5 27019			A 22/02	T. 1011	Home saniens ribosomal protein L13a (RPL13A), mRNA
8952						TOGGGGGGG	AT IT STIRFAMILY J SEQUENCE CONTAMINATION WARNING ENTRY
3508	3 13424	23227	7 3.85	5 6.0E-20	01-39188	SWISSPACE	

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				<u> </u>	? -		
Probe SEQ ID NO:	Exan SEO ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
	1			L	A A 700765 4	HOT HIMAN	ca35b08.s1 NCI_CGAP_GCB1 Homo sapiens cDNA clons IMAGE:1308935 3' similar to contains MEK4.bZ MER4 repetitive element :
8983				2.0E-20	HER371 1	FST HUMAN	CHR220310 Chromosome 22 exon Homo sapieris cDNA clone C22_391 5'
9574	19482	87107	2				Z11d06.r1 NCI_CGAP_GCB1 Homo septens cDNA clone IMAGE:712811 5' similar to contains MER19.72
1987	12648	21751	5.25	1.0E-20	AA281961.1	EST_HUMAN	MEKT9 repende element, how caniens clove IMAGE:3135155 3' similar to contains L1.2 L1
000	l	24018	-	1.0E-20	BF115158.1	EST_HUMAN	hr84bbb.X1 NCI_CCAP_Ind 1 Figure saprens control of the control of
7285	1	L	2.42		-	TN.	Homo sapiens Autosomal Highly Conserved Protein (AHCP), mrNA
3	L						Homo sapiens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, ara pa uei cue, exerimentor
8839	18652	28940	2.81	1.0E-20	AF223391.1	Ę	spliced
							nesogos ri Noi Coar Tri nomo septems como demo mano como como como como como como como co
9321	19011		1.84			EST_HUMAN	repetitive element;
2883	12810		1.08	9.0E-21		EST HOMAN	AJUUSO 14 Selectud Gillusussiin se i oora kinseisi ja maasaa ka ja maa
9045	<u> </u>		2.35	9.0E-21	AW898189.1	EST HUMAN	RCS-NNODBEAUSUSCIO-221-502 INVICED INTO SERVING SERVING SERVING HUMAN
	<u>L</u> _		7.2.7	9 OF 24	AW674891 1	EST HUMAN	bb30e02.71 NIH_MGC_10 Homo septens curry cons invacations of the constant of t
7115	7880	20000		1		EST HUMAN	ob71f06.s1 NCI_CGAP_GCB1 Homo saplens cDNA clone IMAGE:1336835 3
0700	_L	1				SWISSPROT	ATP SYNTHASE A CHAIN (PROTEIN 6)
9207	_ 1			1		CMISCOROT	I AMININ BETA-2 CHAIN PRECURSOR (S-LAMININ) (LAMININ CHAIN B3)
2023	11914				F15000	TOUGH TOUGH	I AMMININ BETA 2 CHAIN PRECURSOR (S-LAMININ) (LAMININ CHAIN B3)
2023	11914	4 21804				SWISSPRO	LAWININ DE TAY CHANGE THE THE THE PROPERTY OF
4181	١.		4.86	7.0E-21	AA046502.1	EST_HUMAN	ZK6 / RUO, T1 Soarles pregnant, ucerus and the company and the
6887	18768	28962	1.43	7.0E-21	AJ277557.1	N	Homo sapiens dNT-2 gene for mitochondrial 5'(3')-deoxyribanucleotidase (dNT-2 gene), exans 1-5
200	1			L	D14718.1	NT	Human chromosomal protein HMG1 related gerie
	L						2g73d03.s1 Soares_fetal_heart_NbHH19W Homo saptens cUNA cione in/ACE.3859501 S String to gar3404338 VITAMIN K-DEPENDENT PROTEIN S PRECURSOR (HUMAN);contains THR.t3 OFR
0808	17971	28220	2.84	t 7.0E-21	1 AA723404.1	EST HUMAN	repetitive element;
S S S	1_			7.0E-21		TN	Homo saplens PTD013 protein (P1D013), mrstyA
4014	L			L	BE40861	EST_HUMAN	601304125F1 NIH_MGC_21 Homo sapiens cuina digne invace. 3030310 3
6	1_			L		5902031 NT	Homo saplens protein tyrosine phosphatase, non-receptor type 21 (F. 15 No.1), intro-
	1		-				om23g03.s1 Soares_NFL_T_GBC_S1 Homo sapiens convaignes invade: 1341900 51111112 3 11 13 12 13 13 13 13 13 13 13 13 13 13 13 13 13
2234	4 12119	9 22021				EST HUMAN	ANTITION OF THE MICE TO HOMO September SONA clone IMAGE:3933880 5
4266	14165				BE96883	ESI HUMAN	Unione smiles and antiden family C. 1 (MAGEC1), mRNA
4698	8 14582	24375	5 5.98	8 5.0E-21	1 · 4885474 NI	N.	DOILO SEPTEMBER LA SETTEMBER LA

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Single Exon Probes Expressed in Heart

Probe SEQ ID NO:	Exen SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Τὸρ Hit Ďescriptor
5109	14977		96.0	5.0E-21	D87675.1	N	Home sapiens DNA for amyloid precursor protein, complets cds
1701	11602	21473	1.24	4.0E-21	AA970713.1	EST HUMAN	0088908.51 NCI_CGAP_Kid5 Homo sepiens cDNA clone IMAGE:1573094 3' similar to TR:Q16530 Q16530 PMS3 MRNA ;contains OFR,t1 OFR repetitive element :
6106	16000	26138	3.06	4.0E-21	AB019576.1	Ν	Rettus norvegicus mRNA for rTIM, complete cds
2228	12113	22015	1.08	3.0E-21	AL163201.2	NT	Homo sapiens chromosome 21 segment HS210001
3041	12968		4.04	3.0E-21	AJ007973.1	NT	Homo sapiens LGMD2B gene
5749			1.72	3.0E-21	BF184739.1	EST_HUMAN	60184465F1 NIH_MGC_54 Homo sapiens cDNA clone IMAGE:4084945 5'
8208			4.79	3.0E-21	BF361093.1	EST_HUMAN	RC1-OT0083-100800-019-g08 OT0083 Homo saplens cDNA
7589			1.55	3.0E-21	AW897760.1	EST HUMAN	CM1-NN0063-280400-203-h08 NN0063 Homo sepiens cDNA
9684		24992	1.32	3.0E-21	AL163213.2	N	Homo sapiens chromosome 21 segment HS210013
140	10114		14.75	2.0E-21	BE163247.1	EST_HUMAN	QV3-HT0458-170200-050-g12 HT0458 Homo sapiens cDNA
919		20688	0.85	2.0E-21	AB007857.2	LN L	Homo saplens mRNA for KIAA0397 protein, partial cds
919					AB007857.2	N	Homo sapiens mRNA for KIAA0397 protein, paritial cds
1196	11106		2.09	2.0E-21	BE064410.1	EST_HUMAN	RC4-BT0311-141199-011-h06 BT0311 Homo capiens cDNA
2599		22361			Q28983	SWISSPROT	ZONADHESIN PRECURSOR
2599	12468		2.89	2.0E-21	Q28983	SWISSPROT	ZONADHESIN PRECURSOR
5370	15200	26126	4,	20 10 0	A 1694509.4	1444	ts30f03.x1 NCI_CGAP_Pan1 Homo sapiens cDNA clone IMAGE:2230109 3' similar to TR:Q99864 Q99854
	L	l			A102+302.1	אואואיסט ופש	THE CITETICAL SIT AD TRUITING
8470	16/49	26944	4.66			EST HUMAN	QV0-HT0103-091199-050-g11 HT0103 Homo sapiens cDNA
77	1		3.43	2.0E-21	AU136779.1	EST_HUMAN	A0135/79 PLACE1 Homo saplens cDNA clone PLACE1005052 5
8412	18287		1.98		BE350127.1	EST HUMAN	ht09g01.x1 NCI_CGAP_Kld13 Homo sapiens cDNA clone IMAGE:3148256.3' similar to contains MER29.b3 MER29 repetitive element:
8636		28778	1.92	2.0E-21	BE973829.1	EST HUMAN	601680836F1 NIH MGC 83 Homo saplens cDNA clone IMAGE:3951008 5
8639	1	28777	1.92		BE973829.1	EST_HUMAN	601680636F1 NIH_MGC_83 Homo saplens cDNA clone IMAGE:3951008 5'
9425	19072		5.13	20E-21	AF176815.1	LN	Homo sapiens putative 8-hydroxyguanine DNA glycosylase gene, complete cds
Š							n/48c04.s1 NCI_CGAP_Pr4 Homo sapiens cDNA clone IMAGE:1043718 similar to contains MER29.b2
227	1	20994	1.68		AA557657.1	EST_HUMAN	MER29 repetitive element;
1381			2.46		AI601264.1	EST_HUMAN	ar88d12.x1 Barstead colon HPLRB7 Homo sapjens cDNA clone IMAGE:2152343 3'
2007	15813		2.43	1.0E-21	AL079752.1	EST_HUMAN	DKFZp434l0830_r1 434 (synonym: https3) Homo sapiens cDNA clone DKFZp434l0830 5
6289	16134	26289	4.43		AI223104.1	EST HUMAN	qg47e05.x1 Soares_testis_NHT Homo:saplens/cDNA clone IMAGE:1838336 3' similar to gb:M64241 QM PROTEIN (HUMAN);
8021	17871		1.46	1.0E-21	5730038 NT	FZ	Homo saplens SET domain and mariner transposase fusion gene (SETMAR) mRNA
4312	14209	23993	2.55	9.0E-22	AI702438.1	EST_HUMAN	1294e03.x1 NCI_CGAP_Kid11 Homo sapiens CDNA clone IMAGE:2286204 3' similar to TR.Q15408 Q15408 NEUTRAL PROTEASE LARGE SUBUNIT

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Top Hit Descriptor	Homo sapiens chromosome 21 segment HS21C001	Homo sapiens chromosome 21 segment HS21 C001	AV781874 MDS Homo sapiens cDNA clone MDSCCG05 5'	AU140358 PLACE2 Homo sapiens cDNA clorie PLACE2000394 5'	Mus musculus TF-1 cell apoptosis related protein-15 (Tfar15), mRNA	CM0-HT0179-281099-076-h05 HT0179 Homo sapiens cDNA	zk67a06.r1 Soares_pregnant_uterus_NbHPU Homo sapiens cDNA clone IMAGE:487858 5'	Homo saplens chromosome 21 segment HS21 C046	ALPHA-2-MACROGLOBULIN PRECURSOR (ALPHA2M)	Homo sapiens gene for activin receptor type IIB, complete cds	Homo sapiens HSPC220 mRNA, complete cds	EST00738 Fetal brain, Stratagene (cat#936203) Homo saplens cDNA clone HFBCF07	Homo sapiens T cell receptor beta locus, TCRBV7S3A2 to TCRBV12S2 region	wx05g07.x1 NCI_CGAP_Gas4 Homo sapiens/cDNA done IMAGE:2542812.3'	Homo sapiens chromosome 21 segment HS21C103	Human dystrophin (DMD) gene, exons 7, 8 and 9, and partial cds	naa27b06.x1 NCI_CGAP_Pr28 Homo sapiens cDNA clone IMAGE:3255898 3' similar to contains Alu	repetitive element;	Homo saplens Xq pseudoautosomal region; segment 1/2	Homo sapiens chromosome 21 segment HS21C002	601882813F1 NIH_MGC_57 Hamo sapiens cDNA clone IMAGE:4095434 5'	Homo saplens chromosome 21 segment HS21C009	tm14h10.x1 NCI_CGAP_Co14 Homo sapiens cDN4 clons IMAGE:2156811 3' similar to gb:L19593 HIGH AFFINITY INTERLEUKIN-8 RECEPTOR B (HUMAN);contains L1.11 L1 repetitive element;	Wi66b04 x1 NCI_CGAP_Brn25 Homo sapiens cDNA done IMAGE:2429839 3' similar to SW:RL21_HUMAN	P46778 60S RIBOSOMAL PROTEIN L21.;	Human chromosomal protein HMG1 related gene	qb28c07.x1 Soares_pregnant_uterus_NbHPU Homo sapiens cDNA clone IMAGE:1697580 3' similar to	contains MER12.t2 MER12 repetitive element;	RC5-BT0707-150300-021-H10 BT0707 Homo sapiens cDNA	yx73d05.s1 Soares melanocyte 2NbHM Homo sapiens cDNA clone IMAGE:267359 3'	IMMEDIATE EARLY GENE 13 PROTEIN PRECURSOR		DAVI-ST0362-261100-001-412-ST0362-Home serious CDNA
Top Hit Database Source	NT	LN	EST_HUMAN	EST_HUMAN	NT	EST_HUMAN	EST_HUMAN	TN	SWISSPROT	۲N	TN	EST_HUMAN	TN	EST_HUMAN	TN	LZ LZ		EST_HUMAN	TN	TN	EST_HUMAN	TN	EST HUMAN		EST_HUMAN	L		EST_HUMAN	EST HUMAN	EST HUMAN	SWISSPROT	TN	LECT LIMAN
Top Hit Acession No.	AL163201.2	AL163201.2	AV761874.1	AU140358.1	_	BE144748.1		AL 163246.2	Q61838	AB008681.1	AF151054.1	M78590.1	AF009660.1	AW029123.1	AL163303.2	U60822.1		BF476511.1	AJ271735.1	AL163202.2	BF218030.1	AL163209.2	AI469679.1		A1859038.1	D14718.1		A1090125.1	BE089841.1	N24942.1	P24916	8394043 NT	1 AMONTON 4
Most Similar (Top) Hit BLAST E Value	9.0E-22	9.0E-22	9.0E-22	9.0E-22	9.0E-22	8.0E-22	8.0E-22	7.0E-22	7.0E-22	7.0E-22	ZZ-30.7	7.0E-22	7.0E-22	6.0E-22	5.0E-22	5.0E-22		5.0E-22	4.0E-22	4.0E-22	4.0E-22	4.0E-22				3.0E-22					2.0E-22	2.0E-22	
Expression Signal	1.26	1.26	3.65	2.92	2.39	4.55	3.8	4.23	2.32	1.18	1.34	2.63	1.86	1.82	2.53	3.97		2.03	0.85	7.2	2.97	1.51	1.2		1.86	1.49		2.72	2.75	2.86	1.33	4.06	ac *
ORF SEQ ID NO:								20401		24613			27599		25955						28244				22288				26864			23091	
Exon SEQ ID NO:	16879	16879	18059	18753	l	10858	16489	10585	14086	14844	16921	17004	17389	16975	15832	17762	1	19213	13498	19767	17995	19315	10868		12397	13534	L			11805	12352	13292	44000
Probe SEQ ID NO:	7002	7002	8171	8945	8997	933	6099	949	4186	4969	7044	7127	7538	9629	5927	7912		9645	3584	6902	8105	9803	943		2523	3820		4695	6793	1910	2476	3373	4133

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Top Hit Descriptor	zo20f01.r1 Soares_senescent_fibroblasts_NbHSF Homo sepiens cDNA done IMAGE:322873 5' similar to gb:X72308 MONOCYTE CHEMOTACTIC PROTEIN 3 PRECURSOR (HUMAN);	RC0-TN0079-150900-025-h12 TN0079 Homo sapiens cDNA	qi76h06.xt Soares_NhHMPu_S1 Homo saplens cDNA clone IMAGE:1878299 3' similar to contains MER29.t3 MER29 repetitive element ;	nv04h11.s1 NCI_CGAP_Pr22 Hamo saplens CDNA clone IMAGE:1219269 3'	nv04h11.s1 NCI_CGAP_Pr22 Homo sapiens cDNA clone IMAGE:1219269 3'	ha24f04.x1 NCI_CGAP_Kid12 Homo saplens cDNA clone IMAGE:2874655 3'	Homo sapiens chromosome 21 segment HS2 (C080	PM4-SN0020-010400-009-h02 SN0020 Homo sapiens cDNA	Human familial Alzheimer's disease (STM2) gene, complete cds	Human DNA, SINE repetitive element	MR0-810659-220200-002-h07 BT0659 Home sapiens cDNA	IL2-UM0076-070400-061-F11 UM0076 Homo sapiens cDNA	Gallus gallus Dach2 protein (Dach2) mRNA, complete cds	HA2340 Human fetal liver cDNA library Homo sapiens cDNA	HA2340 Human fetal liver cDNA library Homo sapiens cDNA	AV647246 GLC Homo sapiens cDNA clone GLCAW C07 3'	Homo sapiens DKFZP56400463 protein (DKFZP56400463), mRNA	Homo saplens Not56 (D. melanogaster)-like protein (NOT56L) mRNA	Rattus norvegicus RIM1B (Rim1B) mRNA, complete cds	Homo saplens chromosome 21 segment HS21 C049	Homo sapiens mannosidase, beta A, lysosomái (MANBA) gene, and ubiquitin-conjugating enzyme E2D 3	(ODESCO) gailes, complete cas Homo seniens mannesidase heta A. Ivensomsi (MANRA) gene and ubloutitin-conjugating enzyme E2D 3	(UBE2D3) genes, complete cds	qg59c03.xf Soeres testis_NHT Homo septens cDNA clone IMAGE:1839460 3' similar to SW:MV10_MOUSE P23249 PROTEIN MOV-10.	Homo sapiens chromosome Xq28 melanoma entigen family A2a (MAGEA2A), melanoma antigen family A12	(MAGEA12), melanoma antigan family A2b (MAGEA2B), melanoma antigan family A3 (MAGEA3), caltractin (CALT), NAD(P)H dehydrogenase-like protein (NSDHL), and Ll>	Pongo pygmaeus olfactory receptor (PPY116) gene, partial cds	Pongo pygmaeus olfactory receptor (PPY116) gene, partial cds
Top Hit Database Source	EST_HUMAN 9	EST_HUMAN R	EST HUMAN M	Т	EST_HUMAN n	EST_HUMAN h		EST_HUMAN P	г		EST_HUMAN N	EST_HUMAN IL	Г		EST_HUMAN H	${f -}$			NT R	I	I		E L	EST_HUMAN S		<u>- 2</u>	NT	NT P
Top Hit Acession No.	N39456.1	2.0E-22 BF092116.1	A1276522.1	AA715315.1	4A715315.1	4W418960.1	4L163280.2	4W865517.1	J50871.1	D14547.1	3E084667.1	AW802801.1	4F198349.1	AI133716.1	AI133716.1	AV647246.1	10092626 NT	5031952 NT	AF199333.1	AL163249.2	7 0007001	4F224009.1	AF224669.1	AI209130.1		U82671.2	AF179818.1	AF179818.1
Most Similar (Top) Hit BLAST E Value	2.0E-22	2.0E-22	2.0E-22		2.0E-22	2.0E-22	2.0E-22	1.0E-22	1.0E-22	1.0E-22	1.0E-22	9.0E-23	8.0E-23	8.0E-23	8.0E-23	7.0E-23	7.0E-23	7.0E-23	6.0E-23	6.0E-23	20.00	0.UE-20.0	6.0E-23	6.0E-23		5.0E-23	5.0E-23	6.0E-23
Expression Signal	1.44	3.39	1.49	7.07	7.07	2.04	1.85	1.78	2.65	1.74	1.49	4.71	0.84	1.48	1.48	1.45	1.04	3.51	1.62	3.12		BR.	1.99	2.18		3.51	3.51	3.04
ORF SEQ ID NO:	25561		27661	L		29071		21609					23238				24438			23846		2000	25339	ļ		25088	25797	
Exan SEQ ID NO:	19447	15655	17448	ı	17490	18779	<u> </u>	ı		ı	16411	ı	13439	15165	L	13192	14649	18274	13305	14071	ŀ	1008	18890			15262		Ш
Probe SEQ ID NO:	5570	5747	7695	7640	7640	8974	6006	1836	2539	3385	6553	8844	3523	5241	6241	3271	4764	8398	3387	4171	2	918	9146	1488		5341	5781	6397

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Top Hit Descriptor	235g09.r1 Soares, pregnant, uterus, NbHPU Homo sapiens cDNA clone IMAGE:503968 5' similar to contains MER29.t2 MER29 repetitive element;	Human endogenous retroviral alament HC2	Human endogenous retroviral element HC2	Homo saplens KIAA0851 gene (partial), XT3 gene and LZTFL1 gene	Human matrix Gla protein (MGP) gene, complete cds		Ė		Т	\neg	v yr16a02.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:205418 67		Homo sapiens cytochrome P450 polypeptide 43 (CYP3A43) gene, partial cds; cytochrome P450 polypeptide	4 (CYP3A4) and cytochrome P450 polypeptide 7 (CYP3A7) genes, complete cds; and cytochrome P450	polypeptide 5 (CYP3A5) gene, partial cds	Human alcohol dehydrogenase gamma subunit (ADH3) gene, exon 1	Homo saplens T cell receptor beta locus, TCRBV7S3A2 to TCRBV12S2 region		Γ	Homo saplens chromosome 21 segment HS21C010	Г	\neg					J QV0-DT0047-170200-122-a06 DT0047 Homo capiens cDNA	Macaca fuscata mRNA for Testis-Specific Protein Y (TSPY), complete cds	Homo sapiens chromosome 21 segment HS21C049	Homo sapiens 959 kb contig between AML1 and CBR1 on chromosome 21q22, segment 3/3	
Top Hit Database Source	EST_HUMAN	۲	ᅜ	Ϋ́	Ν	SWISSPROT	SWISSPROT		EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN			N	NT	FN	EST_HUMAN	N.	NT	EST_HUMAN	EST_HUMAN		EST HUMAN	SWISSPROT	SWISSPROT	EST_HUMAN	NT	N	N F	EST_HUMAN
Top Hit Acesslan No.	AA130165.1	Z70664.1	Z70664.1	AJ289880.1	M55270.1	P22105	P22105		AI201458.1	BE165980.1	H59931.1	H59931.1			AF280107.1	M32658.1	AF009660.1	AU133931.1	AL163252.2	AL163210.2	BE378471.1	AA448097.1		AA683213.1	P23269	P23269	AW937954.1	AB001421.1	AL163249.2	AJ229043.1	AA594178.1
Most Similar (Top) Hit BLAST E Value	3.0E-23	3.0E-23	3.0E-23	2.0E-23	2.0E-23	2.0E-23	2.0E-23		2.0E-23	2.0E-23	2.0E-23	2.0E-23				2.0E-23		2.0E-23	1.0E-23	1.0E-23	1.0E-23	1.0E-23		9.0E-24		8.0E-24	7.0E-24		6.0E-24	6.0E-24	4.0E-24
Expression Signal	3.58	3.61	3.61	4.36	2.77	1.06	1.08	,	1.46	3.8	2.98	2.98			5.62	2.69	2.47	2.02	1.6	4.49	2.91	4.54	,	1.88	1,08	1.08	1.31	2.4	10.14	7.18	3.06
ORF SEQ ID NO:	26653			20402		22519	22520				23589								24110			26937			١	24226			20595	23585	25609
Exon SEQ ID NO:		17233					12627			13569	13804	13804			16475	18888	19218	19676	14323	14543	15937	16744	,	10481	14442	14442	13708	10623	10748	13800	15526
Probe SEQ ID NO:	6580	7329	7329	650	1126	2765	2765	1000	2250	S	3894	3894			6595	9131	9656	9774	4428	4657	6034	6865		240	4549	4549	3796	089	820	3889	5611

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Top Hit Descriptor	Homo sapiens mRNA for KIAA1093 protein, parial cds	Homo sapiens G-2 and S-phase expressed 1 (G13E1), mrivA	hh68c08.x1 NCI_CGAP_GU1 Homo seplens cLNA clone IMACE.2507550 5 similar to contain the contains must be made in the contains the contai	Homo sepiens chromosome 21 segment HS21C052	BOYSHOWSET NIH MGC 46 Hamo sapiens cDNA clone IMAGE:4053396 5'	22/14/10 r1 Stratanene fetal retine 937202 Homo saplens cDNA clone IMAGE:609161 5	2011163;1 Outsing Street Commission (1997) 1 Str	NOS-VANDOUGOSCOCO CONTRACTOR DE LA TRACTOR D	United Comity dispersed repeat element	Human O talling dispersed repeat demonstration of 197 months in RNA	Homo sapiens Col-12/ protein (LOCO10-15), in the sanishe CDNA	V0-S10284-100400-109-010 010284-110110 deptod 5 do 50	Mus megculus mKNA Tor no I rerain, ballari ous	Homo sapiens PTEN (PTEN) gene, exon 2	Homo, sapiens chromosome 21 segment HSZ10103	CMG-NN1010-130300-281-407 NN1010 Homo saptens cours	ne92e10.s1 NC _CGAP_Kid1 Hamo sapiens civins civins living311121 Siling in the same same same same same same same sam	MERT repeditive element.	repotitive element ;	nt25h06.s1 NCI_CGAP_Pr1 Home sapiens cDNA cione IMAGE:914643 Similiar to 3W.1717-7.1701. P36105 PROBABLE 60S RIBOSOMAL PROTEIN L14EA.;	zh65h07,r1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:410989 5	Mus musculus ctogelin (Otog), mRNA	QV2-LT0051-260300-111-f03 LT0051 Homo sapiens cDNA	EST391217 MAGE resequences, MAGP Home saplens cDNA	ye58h04.r1 Soares fetal liver spleen 1NFLS Homo saplens cDNA clone IMAGE:121783 5	PM30T0093-280200-001-g07 OT0093 Homo'sapiens cDNA	OV3-HT0543-140400-149-e11 HT0543 Homo sapiens cDNA	Home saniens hypothetical protein FLJ20344 (FLJ20344), mRNA	House content hundhalted protein Ft J20344 (FLJ20344), mRNA	HOLING SEPTEMBER IN PROPERTY (MALLIKERIN INHIBITOR) (PROTEASE INHIBITOR 4)	NACESTATING PRESENCE 21 Regment HS21C010	אמוועס מקוימונים כיוו מיוויססייים ביים ביים מיוויססייים ביים מיוויססייים ביים ביים ביים ביים ביים ביים בי
Top Hit Database Source	T. L		HUMAN M	T	H CI IMANI	Т	CIMAN	Т	HOMAIN			EST HUMAN	┑			EST_HUMAN C		EST_HUMAN IN	EST_HUMAN	FST HUMAN	П		T HUMAN	Г	Г		Т	NO.			SWISSPROI	7
Top Hit Acession	4.0E-24 AB029016.1	11418318 NT	M814871 1	Ī		1	1	W898189.1	-		8346	1.0E-24 AW820194.1		1.0E-24 AF143313.1		1		7.0E-25 AA483944.1	7.0E-25 AA468646.1	A A 503540 1	W87823.1	7305360 NT	AW8381					HE1/083		8923321	P29622	AL163210.2
Most Similar (Top) Hit BLAST E	4.0E-24	4.0E-24	9 05 34	3.0E-24	3.0E-24	3.0E-24 E	2.0E-24 /	2.0E-24 /	2.0E-24 AI	2.0E-24 M28877.1	1.0E-24	1.0E-24	1.0E-24	1.0E-24	1.0E-24	1.0E-24		7.0E-25	7.0E-25				Ţ	Ĺ	3.0C-25			١				3.0E-25
Expression Signal	2.12	1.53	000	3.02	4.12	5.14	2.33	1.01	3.14	6.55	2.18	1.87	0.91	1.83	4.07	1.88		2.32	3.75		A 32					2.00	2.78			2.66		2.47
ORF SEQ ID NO:	25293	25203			27463	25273	22078		27146		21438		22710		26484			24575	28855		C#067		00207	1		10212				23000		3 26926
Exon SEQ ID NO:	10121	10300		16784	17258	19171	12180	13841	16952	19717	ı	Ł	Į.	1_	1	1	L	14805	18884	1	18750	L	1	\perp				14119	L_	1	<u> </u>	16733
Probe SEQ ID NO:	9080	27/0	2	9069	7449	9587	2298	3729	7075	9433	1670	888	2987	4173	AAEG AAEG	200		4926	9205	3	8942	4/10	6543	2010	8634	1430	3356	4221	3278	3278	4798	6854

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Table 4
Single Exon Probes Expressed in Heart

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						_						_	_			_										
Top Hit Descriptor	as38h08.x1 Barstead aorta HPLRB6 Homo sapiens cDNA clone IMAGE:2319519 3' similar to WP:F49C12.11 CE03371;	as38h08.x1 Barstead aorta HPLRB6 Homo sapiens cDNA clone IMAGE:2319519 3' similar to WP:F49C12.11 CE03371	wg65e08.x1 Soares_NSF_F8_9W_OT_PA_P_S1 Homo saplens cDNA clone IMAGE:2369966 3' similar to contains Alu repetitive element:	EST33446 Embryo, 12 week II Homo sapiens cDNA 5' end	Homo saplens upstream binding transcription frictor. RNA polymerase I (UBTF), mRNA	601191345F1 NIH MGC 7 Homo saplens cDNA clone IMAGE:3535210 5	DKFZp434l066_r1 434 (synonym: htes3) Homo saplens cDNA clone DKFZp434l066 5	2730d08.11 Stratagene neuroepithelium NT2RAMI 937234 Homo sepiens cDNA clone IMAGE:548943 6' similar to gb:M14338 VITAMIN K-DEPENDENT PROTEIN S PRECURSOR (HUMAN).	2630f10.rl Stratagene colon (#837204) Homo capiens cDNA clone IMAGE:588427 5' similar to TR:G695374 G695374 THYROID RECEPTOR INTERACTOR	2030f10.r1 Stratagene colon (#837204) Homo sapiens cDNA clone IMAGE:588427 5' similar to TR:G695374 G895374 THYROID PECEDTOB INTERACTOR	601864963F1 NIH MGC 57 Hamo saciens cDNA clane IMAGE 1083278 st	Homo saplens MLL (MLL) gene, exons 1-3, and partial cds	QV2-PT0012-040400-124-e05 PT0012 Home september aDNA	QV2-PT0012-040400-124-e05 PT0012 Homo sepiens cDNA	nn37d05.s1 NCI_CGAP_GC5 Homo sepiens cDNA clone IMAGE:1086057 3' similar to contains OFR.t1 OFR repetitive element:	Homo sapiens chromosome 21 segment HS210082	DKFZp586L171_s1 586 (symonym: hfkd2) Homo sapiens cDNA clone DKFZp566L171 3'	M.musculus mRNA for astrocytic phosphoprotein, PEA-15	Homo sapiens DNA for amyloid precursor protein, complete cds	to89aD1.x1 NCI_CGAP_Gas4 Homo sapiens cDNA clone IMAGE:2185416 3' similar to contains Alu repetitive element contains Alu	Homo sapiens MHC class 1 region	Homo saplens mRNA for KIAA1438 protein partial cds	QV4-HT0538-020300-123-a02 HT0538 Homo sapiens cDNA	DKFZp434H1910_r1 434 (synonym: https3) Homo saplens cDNA clone DKFZp434H1910 5'	MR2-BN0114-240500-030-907 BN0114 Homo sapiens cDNA	Homo saplens glyceraldehyde-3-phosphate dehydrogenase (GADPH) mRNA, complete cds
Top Hit Database Source	EST_HUMAN	EST_HUMAN	EST HUMAN	EST HUMAN	Ę	EST HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	NAMI H	EST HUMAN	NT	EST HUMAN	EST_HUMAN	EST HUMAN	TN	EST_HUMAN	TN	TN	EST HIMAN	LN	LN L	EST_HUMAN	EST_HUMAN	T_HUMAN	NT
Top Hit Acessian No.	AI708235.1	AI708235.1	AI761429.1	AA329548.1	7857670 NT	BE266187.1.	AL045855.2	AA115885.1	AA152464.1	AA152484 1	BF245458.1	AF038405.1	AW875651.1	AW875651.1	AA583173.1	AL163282.2	AL038089.2	X86694.1	D87675.1	A[801412.1	AF055066.1	AB037859.1	BE170371.1	AL039363.2	BE814995.1	AF261085.1
Most Similar (Top) Hit BLAST E Value	5.0E-28	5.0E-26	5.0E-26	4.0E-28	4.0E-26	4.0E-28	3.0E-26	3.0E-26	3.0E-26	_			_	3.0E-26	3.0E-26	2.0E-26		-	2.0E-26	2.0E-26		_	1.0E-26	1.0E-26	_	1.0E-28 /
Expression Signal	3.33	3.33	1.71	1.52	3.77	3.74	1.5	2.41	1.19	1.19	4.35	2.18	1.99	1.99	10.55	5.61	3.36	4.28	2.88	4.55	1.82	2.19	2.36	1.39	0.84	16.79
ORF SEQ ID NO:	20918	20919				28194	21740		23408	23409	26114		28951	28952	28984	20418		22924		28690		-	18927	21786	22282	
Exon SEQ ID NO:	11073	11073	19435		17283	17944	11853	11880	13625	13625	15978	17998	18665	18665	18691	10800	11722	13118	18023	18420	18537	18962	10106	11894	12380	12521
Probe SEQ ID NO:	1160	1160	8976	1525	7416	8053	1958	1987	3712	3712	6131	8108	8853	8853	8879	999	1826	3183	8135	8549	8720	9252	132	2001	2516	2654

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Single Exon Probes Expressed in Heart

	_	-	_	_	_										_						_			_		
Tcp Hit Descriptor	MR3-HT0487-150200-113-g01 HT0487 Homo saplens cDNA	DKFZp566C2146_r1 568 (synonym: hfkd2) Homo saplens cDNA clone DKFZp566C2146 5	CHR220032 Chromosome 22 exan Homo saplens cDNA clane C22_45 5'	Homo sapiens MAGE-B2 (MAGE-B2), MAGE-E3 (MAGE-B3), MAGE-B4 (MAGE-B4), and MAGE-B1	(WASE-51) genes, complete cas	MYOMESIN 2 (M-PROTEIN) (165 KD TITIN-ASSOCIATED PROTEIN) (165 KD CONNECTIN- ASSOCIATED PROTEIN)	naa03c07 xt NCI_CGAP_Pr28 Homo sapiens cDNA clone IMAGE:3253644 3' similar to contains OFR.t1 OFR repatitive element;	Wi49c04.x1 NCI_CGAP_Lu19 Homo sepiens cDNA done IMAGE:2406150 3' similar to contains THR.b2 THR repetitive element;	Homo saplens chromosome 21 segment HS21C027	au87h08.x1 Schneider fetal brain 00004 Homo sapiens cDNA clone IMAGE:2783295 3' similar to gb:K00558 TUBULIN ALPHA-1 CHAIN (HUMAN);	au87h08.x1 Schneider fetal brain 00004 Homo sapiens cDNA clone IMAGE:2783295 3' similar to gb:K00558 TUBULIN ALPHA-1 CHAIN (HUMAN):	PM2-SN0018-220300-002-a07 SN0018 Homo sapiens cDNA	ADP,ATP CARRIER PROTEIN, LIVER ISOFORM T2 (ADP/ATP TRANSLOCASE 3) (ADENINE NUCLEOTIDE TRANSLOCATOR 3) (ANT 3)	Homo sapians WRN (WRN) gene, complete cds	MR4-BT0398-250800-204-d06 BT0398 Homo sapiens cDNA	J1751F Human fetal heart, Lambda ZAP Express Homo sapiens cDNA clone J1751 5' similar to REPETITIVE ELEMENT L1	CM1-CT0315-091299-063-407 CT0315 Hamo saplens cDNA	CM1-CT0315-091299-063-407 CT0315 Homo sapiens cDNA	Human endogenous retroviral element HC2	hi51h12.x1 Soares, NFL_T_GBC_S1 Homo squens cDNA clone IMAGE:2978879 3' similar to TR:O76040 O76040 ORF2: FUNCTION UNKNOWN.	Homo saplens Xq pseudoautosomal region; segment 1/2	AV723365 HTB Homo sapiens cDNA clone HTBAHE02 5'	Human nucleolar protein (B23) mRNA, complete cds	602121491F1 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:4278527 6'	602121491F1 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:4278527 5	Mus musculus sperm fall associated protein (Stap), mRNA
Top Hit Database Source	EST_HUMAN	EST_HUMAN	EST_HUMAN	<u> </u>	Z	SWISSPROT	EST_HUMAN	EST HUMAN	Ę	EST_HUMAN	EST HUMAN	EST_HUMAN	SWISSPROT	N	EST_HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	F	EST HUMAN	F	EST_HUMAN	TN	EST_HUMAN	EST_HUMAN	ᅜ
Top Hit Acession No.	BE165980.1	AL038487.1	H55083.1		บรราชร.า	P54296	BF445556.1	A1831462.1	AL 163227.2	AW162737.1	AW162737.1	AW864776.1	; P12236	AF181897.1	BĘ926560.1		AW857579.1	AW857579.1	Z70664.1	AW629172.1	AJ271735.1	AV723365.1		BF666614.1	BF666614.1	9910569 NT
Most Similar (Top) Hit BLAST E Value	1.0E-26	1.0E-26	1.0E-26	100	8.0E-2/	9.0E-27	9.0E-27	8.0E-27	8.0E-27	8.0E-27	8.0E-27	8.0E-27	8.0E-27	8.0E-27	8.0E-27	8.0E-27	8.0E-27	8.0E-27	7.0E-27	7.0E-27	7.0E-27	7.0E-27	6.0E-27	5.0E-27	5.0E-27	4.0E-27
Expression Signal	2.75	3.17	1.84	0	6.11	3.48	3.97	3.09	4.13	18.87	18.87	98.0	3.31	0.91	3.12	4	1.68	1.68	1.22	2.09	4.22	2.07			2.82	1.54
ORF SEQ ID NO:						28118		19787		21168	21159		22874	23035		26088	27386									26080
Exon SEQ ID NO:			19737		740/-	19470	18812	9666	10486	11300	11300	12009	13073	13230	15122	15956	17186	17186	10602	14896	18020	19204	17999	17727		15949
Probe SEQ ID NO:	6089	8266	9493	7987	1304	8027	9013	10	545	1395	1395	2121	3148	3309	6165	6196	7310	7310	899	5023	8132	9631	8109	7877	7877	6046

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Single Exon Probes Expressed in Heart

					_	_							_														
Top Hit Descriptor	Raftus norvegicus putative four repeat lon channel mRNA complete cds	H.sapiens DNA for endogenous retroviral like elament	Rrattus RYA3 mRNA for a potential ligand-binding protein	PM0-BT0527-090100-001-d11 BT0527 Home seniens cDNA	7844C08 Chromosome 7 Fetal Brain cDNA I Ibrary Home saniens cDNA clare 2044C08	601458531F1 NIM MGC 68 Home seniens CDNA clone MARGE-38820888 F	Homo sapiens alpha NAC mRNA, complete cds.	nk01b10.s1 NCI_CGAP_Pr11 Homo sapiens cDNA clone IMAGE:1000699 similar to gb:M17886 60S ACIDIC RIBOSOMAL PROTEIN P1 /HIJMAN	hI51h12.x1 Scares_NFL_T_GBC_S1 Homo sapilens cDNA clone IMAGE:2975879 3' similar to TR:O76040 O76040 ORF2: FUNCTION UNKNOWN ;	Homo saplens jun dimerization protein gene, parital cds; cfos gene, complete cds: and unknown gene	Homo saplens fun dimerization profein nene narital crie: rine nana comolate ode: ond universe	Rattus noviecicus voltace-cated sodium channel mRNA complete cus, enu unixnown gene	W/28g07,x1 NCI CGAP Utf Home seniens cDNA clone IMACE-2228288 2	nh08h05.s1 NCI_CGAP_Thy1 Homo sapiens cDNA clone IMAGE:943737 similar to contains L1.t3 L1 repetitive element	EST00738 Fetal brain Stratagene (cattloge208) Home conject CDNA class LIED OF 2	EST00738 Fetal brain. Stratagene (cattle336206) Homo seniens cDNA clone LED CE07	AU121685 MAMMA1 Homo saplens cDNA clone MAMMA1000746 5	Inko1b10.s1 NCI_CGAP_PT11 Homo saplens clone IMAGE:1000699 similar to gb:M17886 60S ACIDIC RIBOSOMAL PROTEIN P1 (HUMAN)	Homo saplens chromosome 21 segment HS21C046	Homo saplens DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes, complete cds)	Homo sapiens Retina-derived POU-domain factor-1 (RPE-1) mRNA	HSPD20461 HM3 Homo sapiens cDNA clone safonnoscoto	HSPD20481 HM3 Homo sapiens cDNA clone s4000095C10	Homo saplens mRNA for KIAA0454 protein, partial cds	RC6-BT0627-140200-011-E06 BT0827 Hamp sapiens cDNA	Human mRNA for KIAA0260 gene, partial cds	Bos taurus latrophilin 3 splice variant bbah mRNA, complete cds
Top Hit Database Source	N	N	N	EST HUMAN	EST HUMAN	Г	Π	EST HUMAN	EST_HUMAN	NT	IN		T HUMAN	T	1	EST HUMAN	1		Т	Į.		T HUMAN	1	ŁN	Т	- LN	LN H
Top Hit Acession No.	AF078779.1	X89211.1	X60658.1	BE071924.1	AA077705.1	BF035327.1	AF054187.1	AA665345.1	AW629172.1	4F111167.2	4F111167.2	AF000368.1	41866347.1	AA551527.1			AU121685.1	2.0E-27 AA565345.1	Г	1.0E-27 AB026898.1	5855			1.0E-27 AB007923.1	Ψ.		1.0E-27 AF111093.1
Most Similer (Top) Hit BLAST E Vælue	4.0E-27	4.0E-27	3.0E-27	3.0E-27	3.0E-27	3.0E-27	2.0E-27	2.0E-27	2.0E-27	2.0E-27	2.0E-27	2.0E-27	2.0E-27	2.0E-27	2.0E-27	2.0E-27	2.0E-27	2.0E-27	1.0E-27	1.0E-27	1.0E-27	1.0E-27 F30158.1	1.0E-27 F30158.1	1.0E-27	1.0E-27	1.0E-27 D87449.1	1.0E-27
Expression Signal	1.23	2.56	6.42	1.27	5.13	2.98	7.86	18.58	10.27	1.43	1.43	1.09	1.5	23	1.22	1.22	2.89	20.82	1.58	1.41	6.31	2.2	2.2	1.72	1.89	2.68	3.14
ORF SEQ ID NO:		28985	21782	23849	24976	27549	19820			22816	22917	23627	26780		27847	27848	28450			20749	25972	26136	28137	27075		27672	28046
SEQ ID			11889	14074	15200	17343	10023	11749	12998	13112	13112	13852	16592	17208	17618	17618	18201	11749	10374	10803	15848	15999	15999	16882	17071	17457	18751
Probe SEQ ID NO:	9999	888	1995	4174	5278	7365	စ္တ	1853	3071	3187	. 3187	3944	8712	7338	7768	7768	8324	8778	429	980	5943	6105	6105	7005	7194	7606	8943

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Single Exon Probes Expressed in Heart

Probe SEQ ID NO:	Exon SEQ (D NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acesslon No.	Top Hit Database Source	Top Hit Descriptor
138	10109		2.32		9.0E-28 BE348399.1	EST HUMAN	hw17c11.x1 NCI_CGAP_Lu24 Homo sepiens oDNA clone IMAGE:3183188 3's lmillar to TR:007314 Q07314 SECRETED NEUREXIN III.44 PHA.C PRECI IRSOR 131 TB-0A7364 TB-0A7344
308	10270	20089			9.0E-28 AU126260.1	EST_HUMAN	AU126260 NT2RP1 Homo sapiens cDNA clone NT2RP1000443 5'
5118	14988	24760	1.21	905-28	9 0E-28 A1590115 1	NAM I	to12b09x1 NCI_CGAP_Utz Homo sepiens cDNA clone IMAGE:2178809 3' similar to contains OFR.t1 OFR
3							to 12b09.x1 NCI_CGAP_U2 Homo sapiens cDNA done IMAGE:2178809 3' similar to contains OFR.t1 OFR
8116		24761	1.21	9.0E-28	9.0E-28 AI590115.1	EST_HUMAN	repetitive element;
808	18866		3.74	9.0E-28	9.0E-28 BF377859.1	EST_HUMAN	CM2-TN0140-070900-372-g01 TN0140 Homo sapiens cDNA
9418	19592		1.97	8.0E-28	8.0E-28 AW157571.1	EST HUMAN	au83h08.x1 Schneider fetal brain 00004 Homo sapiens cDNA clone IMAGE:2782911 3' similar to
1164	11077	20922	7.54	7.0E-28 AU	142750.1	EST HUMAN	AU142750 Y79AA1 Home sapiens of NA close Y79A44nona24 5
8523	18395	28660	2.54	7.0E-28	11417866	1	Homo sapiens gamma-diutamyfransferase_like/a-thyty 1 (CCT) 41\ mbni A
9053	18836		1.44	7.0E-28	7.0E-28 AV735348.1	EST HUMAN	AV735348 CB Homo sepiens cDNA clone CREAKA12 5
3987			1.27	6.0E-28			Homo sapiens mRNA for KIAA0866 nortein complete case
3987	13894	23671	1.27	6.0E-28	6.0E-28 AB020673.1	LN	Homo sapiens mRNA for KIA40866 protein complete cds
9673	19232		2.73	805.38	8 OF 28 A B G 4 5 5 3 1	ECT LILIAM	aa60e03.r1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:825340 5' similar to contains Alu
	L			22700	T. SOLLOW	NEW TOWN	rependive element contains element PTK6 repetitive element;
315	10277		3.08	5.0E-28		EST HUMAN	wo18c07.x1 NCI_CGAP_Pan1 Homo saplens cDNA clone IMAGE:2455692.3' sImilar to contains THR.b1 THR repetitive element:
3927	13836	23616	1.85	5.0E-28	5.0E-28 R79762.1	EST_HUMAN	y89f10.r1 Soares placenta NbZHP Homo sapiens cDNA clone IMAGE:146443 5'
2584	12455		1.42	4.0E-28	4.0E-28 AW195066.1	EST HUMAN	xn33c09.x1 NCI_CGAP_Kid11 Homo saplens cDNA clone IMAGE:2695504 3' similar to SW:GG95_HUMAN CO8379 GOI GIN.95
3070	12997	22788	3.39	4.0E-28		EST_HUMAN	601300703F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3636305 5
6336	16199	26359	1.59	4.0E-28	4.0E-28 AI198941.1	EST HUMAN	466010.x1 Sogres_tests_NHT Homo sapiens CDNA clone IMAGE:1755019 3' similar to gb:M19503 LINE-1 REVERSE TRANSCRIPTASE HOMOI OG (HI IMAN):
8230	18140		ξ	10 10 7			
8371			53.6	4.0E-28/	38241 1	Z	name sapiens chromosome 9 duplication of the T cell receptor beta locus and trypsinogen gene families
							die Genes Car Dri min'n für gijverlauden yde-ki-phosphate denydrogenase, complete das die Genes festis NHT Homo saniens china MASE 17755010 of circulation in the contraction of circulations and contraction in the contraction of circulations and contraction in the contraction of circulations and contraction of circulations and contraction of circulations and contraction of circulations and contraction of contractions and contraction of contractions and contraction of contractions and contraction of contractions and contractions are contracted and contractions and contractions and contractions and contractions and contractions and contractions and contractions are contracted and contractions and contractions and contractions are contracted and contractions and contractions are contracted and contractions and contractions are contracted and contractions and contractions are contracted and contractions and contractions are contracted and contractions and contractions are contracted and contractions and contractions are contracted and contracted and contractions are contracted and contracted and contractions are contracted and cont
8388	16199	26359	2.94	4.0E-28	4.0E-28 AI198941.1	EST_HUMAN	REVERSE TRANSCRIPTASE HOMOLOG (HUMAN);
1262	11169		1.89	3.0E-28 AF1	55382.1	LN.	Homo saplens metalloprotease-like, disintegrin-like, cysteine-rich protein 2 epsilon (ADAM22) mRNA, complete eds
7124	17001	27183	2.19	3.0E-28 BF3	54030.1		MR3-HT0713-280500-013-f09 HT0713 Homo similans cDNA
8306	18183	28430	1.84	3.0E-28 U53	588.1	LN L	Homo sapiens MHC class 1 region
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Single Exon Probes Expressed in Heart

Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
9489	19106		2.44	3.0E-28	A1831991.1	EST HUMAN	wj98f07.x1 NCI_CGAP_Lym12 Homo sapiens CDNA clone IMAGE:2410885 3' similar to contains Alu repetitive element contains element HGR repetitive element
88			6.45	2.0E-28	2.0E-28 BE062167.1	EST HUMAN	RC1-BT0254-220300-019-c05 BT0254 Homoleaplens cDNA
1149	11062	20905	10	2.0E-28	2.0E-28 Y11107.3	FZ	Homo saplens ITGB4 gene for integrin beta 4 subunit, exons 3-41
2427	12304	22200	,	A 90 30 0	3496344	TOD TOD	qo35b06.x1 NCI_CGAP_Lu5 Homo sepiens cDNA clone IMAGE:1910483 3' similar to contains L1.b2 L1
5821			4.28	2.0E-28		EST HIMAN	FORBITATORET NIH MGC 54 Home conjune con 1200 NA CELLA MACE AND MGC 54 Home conjune con 1200 NA CELLA MACE 21
7528	L. I		5.88	2.0E-28	2.0E-28 AW972305.1	EST HUMAN	EST384394 MAGE resequences. MAGL Homo septems cDNA
1000							Homo sapiens mannosidase, beta A, Iysosomal (MANBA) gene, and ubiquitin-conjugating enzyme E2D 3
288	- 1	28992	227	2.0E-28 AF		۲	(UBE2D3) genes, complete cds
1463		21233	2.42	1.0E-28	1.0E-28 D38044.1		Human gene for Ah-receptor, exon 7-9
2173		21963	1.55	1.0E-28	BF333236.1	EST_HUMAN	QV1-BT0821-120900-360-b03 BT0821 Homo saptens cDNA
2650		22407	1.03	1.0E-28	1.0E-28 AF000995.1	ΡΉ	Homo saplens ubiquitous TPR motif, Y isoform (UTY) mRNA, alternative transcript 2, complete cds
6587	ı		4.48	1.0E-28	11429885 NT	占	Homo saplens similar to ribosomal protein L12 (H. sapiens) (LOC63091), mRNA
6875	16555		3.02	1.0E-28	8922793 NT	F	Homo sapiens hypothetical protein FLJ10968 (FLJ10968), mRNA
7346	17214	27413	2 63	1 0F-28	A & 3/18744 4	INOMI IO TOD	EST179615 HCC cell line (matastasis to liver in mouse) Il Homo sapiens cDNA 6' end cimilar to similar to
7678	1	27753	5.15	1 OF 28	4758434 INT		Homo contain calliforni floranciana illocation and Corresponding
7678	L	27754	5.15	1 OF 28	4758434 NT		The capture of sections of the capture of the captu
8028	L		3.95	1.0E-28	1.0E-28 AA054182 1	EST HIMAN	753-01 1 Soares cathe NOVAHD Home control of the 1140 From 100 Fro
9797	19485		1.43	1.0E-28	T		Homo sapiens chromosome 21 segment NS21/047
2686	19691	24896	2.8	9.0E-29		EST HUMAN	h/76q06.x1 Soares NFL T GBC S1 Homp saniens CDNA Appe M/4 CE-2021228 2
9585	19169		2:92	8.0E-29 Q00130		SWISSPROT	HYPOTHETICAL GENE 50 PROTEIN
1584	11488	21349	0.91	7.0E-29	N966447.1	EST_HUMAN	EST378521 MAGE resequences, MAGI Homolsapiens cDNA
9942	19410		5.08	7.0E-29 A.	1132352.1	NT	Rattus norvegicus mRNA for 45 kDa secretory protein, partial
Car	0.4	30000	q	. 0			wp69b01.x1 NCL_CGAP_Bm25 Homo sapiens cDNA clone IMAGE:2466985 3' similar to TR:015475
9353	19028	C7507	0.02 2 88	9.0E-29	0.0E-29 AI936/48.1	FOT HUMAN	U154/5 UNNAMED HERV-H PROTEIN ;contains LTR7.51 LTR7 repetitive element;
		1	99.5	0.00-23	1	╗	KC3-C I UV6Z-Z U8UU-UZ1-c05 U I 006Z Homo sapiens cDNA
24.8 25.8	190/8		1.86	6.0E-29		T HUMAN	602184092F1 NIH_MGC_42 Homo saplens cDNA clone IMAGE:4300079 5'
4928	14807		0.98	5.0E-29,	5.0E-29 AL163203.2		Homo saplens chromosome 21 segment HS210003
7068	16945		7.61	5.0E-29,		EST_HUMAN	RC3-OT0091-170300-011-c12 OT0091 Homo saplens cDNA
3194	13119		1.68	4.0E-29/		EST_HUMAN	cn15c02.x1 Normal Human Trabecular Bone Cells Homo sapiens cDNA clone NHTRC 2015-02 random
5858	15568	-	6.65	4.0E-291	4.0E-29 BE164930.1		QV1-HT0471-280300-121-a05 HT0471 Hamo saniens cDNA
						ı	Large granders arranged

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. Top Hit Descriptor	Human 90 kD heat shock protein dene, complete ods	Homo saplens PTS gene for 6-pyruvovitetrahydronterin synthese complete cds	QV1-BT0821-120900-360-b03 BT0821 Homo sapiens cDNA	Human gene for Ah-receptor, exon 7-9	xv17f03.xt Soares_NFL_T_GBC_S1 Home sepiens cDNA clone IMAGE:2813405 3' similar to contains Alurement contains MEB10 49 MEB10 49 MEB10 40	Homo sapiens chromosome 21 segment HS21(2046	262b01.r1 Soares_lestis_NHT Homo sapiens cDNA clone IMAGE:726889 5' similar to TR:G1335769	Human Hs IM15 mRNA for Hs Im15 complete ode	Homo sepiens envelone protein RICA (any) gains complete ade	Homo saplens envelope protein RIC-6 (env) dene, complete cds	wr65d10.x1 NCI_CGAP_UTI Home sapiens cDNA clone IMAGE:2492563 3' similar to TR:016646 015546 HERV-E ENVELOPE GLYCOPROTEIN:	wr85d10.x1 NCI_CGAP_Uf1 Homo sapiens cDNA clone IMAGE;2492563 3' sImilar to TR:O15546 O16548 HERV-E ENVELOPE GI YCOPROTFIN	Horno saplens chromosome 21 segment HS210068	wt27g07.xf Soares_NF_T_GBC_S1 Homo sepiens cDNA clone IMAGE:2356860 3' similar to contains element MER6 repetitive element :	wt27g07.x1 Soares_NFL_T_GBC_S1 Homo sépiens cDNA clone IMAGE:2356860 3' similar to contains element MFR6 renefitive element :	Homo saplens chromosome 21 segment HS21(2)48	Homo saplens chromosome 21 segment HS21C048	Homo sapiens chromosome 21 segment HS21(3048	Homo sapiens chromosome 21 segment HS210048	QV0-OT0032-080300-155-d01 OT0032 Homo saplens cDNA	Homo sapiens chromosome 21 segment HS21C027	RC1-HN0003-220300-021-b04 HN0003 Homo/saplens cDNA	nz20c07.s1 NCI_CGAP_GCB1 Homo saplens cDNA clone IMAGE:1288332 3' similar to contains MER4.b1	WICK+ repeare gernent;	Homo sapiens zinciiron regulated transporter-like (ZIRTL), mRNA	HSC23F051 normalized infant brain cDNA Homo sapiens cDNA clone c-23f05	ESTS/31/ Inymus I Homo sepiens cDNA 5 end similar to EST containing O family repeat	F12.1 3 B11.1 tumorz Homo sapiens cDNA 3
Top Hit Database Source	TN	TN	EST HUMAN .	TN	FST HIMAN	LN	FST HIMAN	L	LX.	TN	EST HUMAN	EST HUMAN	NT	EST_HUMAN	EST HUMAN	N I	FN	Į.	F	EST_HUMAN	LZ	EST_HUMAN		NAMOL		Т	Т	ESI_HUMAN
Top Hit Acession Na.	J04988.1	AB042297.1	BF333236.1	D38044.1	4W303317.1	AL163246.2	AA403053 1	J63882 1	AF084869.1	AF084869.1	A1963604.1	1963604.1		AI806418.1	AI806418.1								0 05 30 0 724245 4	27.70	C\$/7741	,	557072 4	23/0/2.1
Most Similar (Top) Hit BLAST E Value	4.0E-29	3.0E-29		3.0E-29	3.0E-29	3.0E-29	3.0E-29	3.0E-29	2.0E-29/	2.0E-29 /	2.0E-29	2.0E-29	2.0E-29	2.0E-29	2.0E-29	2.0E-29	2.0E-29	2.0E-29	2.0E-29	2.0E-29	2.0E-29 A	1.0E-29	00 00	9.00.0	8.05.30	8.0E-30 F08688.1	8.0E-30.A	סיטביטטים
Expression Signal	4.71	1.45	1.31	2.07	9,1	1.64	2.22	1.51	1.43	1.43	6.12	6.12	2.03	1.43	1.43	2.95	2.95	3.39	3.39	1.96	1.73	5.41	2	,	0.1	8.81	3.53	122.2
ORF SEQ ID NO:	27149	23997	24305	27138	27429		28728		20240	20241	21278	21279	23855	25765	25765	27539	27540	27973	27974			27174	25005		1	2880	27105	1221 12
Exon SEQ ID NO:	16956			16947	17228	17362	18459	18959	10427	10427	11421	11421	14082	15658	15658	17334	17334	17729	17729	18618	18807	16982	15874	18880	16734	16691	16916	7.1
Probe SEQ ID NO:	2079	4317	4628	7070	7362	7492	8591	9248	484	484	1516	1516	4182	5750	6461	7474	7474	7879	7879	8804	9004	7106	5968	9133	5948	6812	7039	

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Single Exon Probes Expressed in Heart

Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
1499	11403		1.16	7.0E-30	7.0E-30 BE091133.1	EST_HUMAN	PM4-BT0724-150400-004-d11 BT0724 Homo sapiens cDNA
1549	11454		0.95	6.0E-30	X51755.1	NT	Human lambda-immunoglobulin constant region complex (germline)
1738	11637	21505	1.26	6.0E-30	D25303.1	NT	Human mRNA for Integrin alpha subunit, complete cds
3153	13078		2.41	8.0E-30	8.0E-30 BE008028.1	EST_HUMAN	QV0-BN0147-290400-214-f12 BN0147 Homo sapiens cDNA
8915	11454		3.15	6.0E-30	X51755.1	NT	Human lambda-Immunoglobulin constant region complex (germline)
			3	i d	7 0000001	i i	tg92g03.x1 NCI_CGAP_CLL1 Homo sapiens cDNA clone IMAGE:2116276 3' similar to contains Alu
3931	13840	73970	31.23	1	5.0E-30 AI398992.1	ES COMAN	Imperior granitate hadraface (ACOO) nene avan 7
2020	1	1	500		007931.1	1	Harrison and instruction of Learners HEDRICATE
8261		00000	3.66	6.0E-30	6.0E-30 AL163278.2	Z	Homo sapiens arromosome 21 segment no.21,0076
8495		28832	9,68	5.0E-30	5.0E-30 AL163Z10.Z	Z	nomo sapiens circinosonia et i segment nomo sapiens como
8495	_]	28633	8.68	5.0E-30	AL163210.2	Ŋ	Homo sapiens chromosome Z1 segment HSZ1CO10
2098		21880	1.64	4.0E-30	4.0E-30 AW937471.1	EST_HUMAN	QV3-DT0043-090200-080-c06 DT0043 Homo saplens cDNA
2098	11985		1.64	4.0E-30	AW937471.1	EST_HUMAN	QV3-DT0043-090200-080-c08 DT0043 Homo saplens cDNA
7162	17039	27231	1.49	4.0E-30	4.0E-30 AW812488.1	EST_HUMAN	CM1-ST0181-091199-035-f08 ST0181 Homo saplens cDNA
							qq93c05x1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA clone IMAGE:1938920 3' similar to
1134	11048		1.71	3.0E-30 AI	Al338551.1	EST_HUMAN	contains MER29.b2 MER29 repetitive element;
3697	13611	23395	0.91	3.0E-30	3.0E-30 AF128893.1	NT	Homo sapiens telomerase reverse transcriptase (TERT) gene, exons 1-6
8538	18410	28675	2.38	3.0E-30	P34056	SWISSPROT	TRANSCRIPTION FACTOR AP-2
629	10593	20411	1.18	2.0E-30	AW857315.1	EST_HUMAN	CM0-CT0307-310100-158-h03 CT0307 Home sapiens cDNA
1068	10984		2:32	2.0E-30	2.0E-30 F08688.1	EST_HUMAN	HSC23F051 normalized infant brain cDNA Homo sapiens cDNA clone o-23f05
1464	11369	21234	5.91	2.0E-30	BE175877.1	EST_HUMAN	RC5-HT0582-110400-013-H08 HT0582 Hamp sapiens cDNA
2683	12548	22438	10.97	2.0E-30	2.0E-30 BE785232.1	EST_HUMAN	IL2-NT0101-280700-116-E04 NT0101 Homo sapiens cDNA
2889	12816		8:38	2.0E-30	AF114156.1	NT	Homo sapiens Y-linked zinc finger protein (ZFY) gene, complete cds
3721	13633	23419	. 2.11	2.0E-30	AW 206581.1	EST_HUMAN	UI-H-BI1-afo-c-12-0-UI.s1 NCI_CGAP_Sub3 Homo saplens cDNA clone IMAGE:2722558 3'
4668	14554	24346	1.72	2.0E-30	BE298945.1	EST_HUMAN	601119860F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:3029438 5'
4668	14554	24347	1.72	2.0E-30	BE298945.1	EST_HUMAN	601119860F1 NIH_MGC_17 Homo septens cDNA clone IMAGE:3029438 5'
9969	16844	27036	3,46	2.0E-30	2.0E-30 C18939.1	EST_HUMAN	C18939 Human placenta cDNA (TFujiwara) Homo saplens cDNA clone GEN-570C01 5'
7019	16896	27085	1.55	2.05-30 8	BE670617.1	EST HUMAN	7e37c12x1 NCI_CGAP_Lu24 Homo sepiens cDNA clone IMAGE:3284662 3' similar to SW:DHSA_HUMAN P31040 SUCCINATE DEHYDROGENASE [ÜBIQUINONE] FLAVOPROTEIN SUBUNIT PRECURSOR;
						1	
7019	16896		1.55	2.0E-30	BE670617.1	EST_HUMAN	7e37e12x1 NCI_CGAP_Lu24 Homo septens oDNA clone IMAGE:3284682 3' similar to SW:DHSA_HUMAN P31e40 SUCCINATE DEHYDROGENASE [JBIQUINONE] FLAVOPROTEIN SUBUNIT PRECURSOR;
7743	i I	27814		2.0E-30	2.0E-30 AW971568.1	EST_HUMAN	EST383657 MAGE resequences, MAGL Honio saplens cDNA

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Probe SEQ ID S NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
7802	17652	27889	4.47	2.0E-30	2.0E-30 AW470791.1	EST_HUMAN	ha33d08.x1 NCI_CGAP_Kid12 Homo sapiens cDNA clone IMAGE:2875499 3' similar to contains THR.b3 THR repetitive element ;
284	10249	20069	10.22	1.0E-30	1.0E-30 C18939.1	EST_HUMAN	C18939 Human placenta cDNA (TFujlwara) Homo sapiens cDNA clone GEN-570C01 5'
527	10469	20281	1.6	1.0E-30	1.0E-30 AW468897.1	EST_HUMAN	hd30b04.x1 Soares_NFL_T_GBC_S1 Homo seplens cDNA clone IMAGE:2910991 3' similar to contains MER1.t3 MER1 MER1 repetitive element ;
669	10632	20457	2.92	1.0E-30	1.0E-30 AL163203.2	TN	Homo sapiens chromosome 21 segment HS21C003
2165	12052	21953	3.24		1.0E-30 AA664377.1	EST-HUMAN	ac77b08.s1 Stratagene lung (#937210) Homo sapiens cDNA clone IMAGE:868599 3'
2413	12290	22187	2.57		1.0E-30 BF347728.1	EST_HUMAN	602022560F1 NCI_CGAP_Brn67 Homo sapleris cDNA clone IMAGE:4157991 5'
2973	12900	22699	1.03	1.0E-30	TN 1605083	NT	Homo saplens methionine aminopeptidase; eIF-2-associated p67 (MNPEP), mRNA
3015	12943	22738	0.93	1.0E-30 A	AA315045.1	EST_HUMAN	EST186868 HCC cell line (matastasis to liver in mouse) II Homo saplens cDNA 5' end
6545	16403	26582	2:32	1.0E-30	1.0E-30 BF183230.1	EST_HUMAN	601809932F1 NIH_MGC_18 Homo sapiens cDNA clone IMAGE:4040694 6'
9733	19615		5.1	1.0E-30		EST_HUMAN	CHR220532 Chromosome 22 exon Homo sapiens cDNA clone C22_728 5'
7013	16890		2.04	9.0E-31		EST_HUMAN	HSC05F032 normalized infant brain cDNA Homo sapiens cDNA clone c-05f03 3'
1060	10977	20820	1.92	8.0E-31	8923388	N L	Homo sapiens hypothetical protein FLJ20420 (FLJ20420), mRNA
2362	12242		5.14	8.0E-31	8.0E-31 AL163208.2	IN	Homo saplens chromosome 21 segment HS21C008
694	10627		1.72	7.0E-31	7.0E-31 AA372637.1	EST_HUMAN	EST84555 Calon adenocarcinoma IV Homo saplens cDNA 5' end
2633	12500	23392	1.7	7.0E-31	7.0E-31 BE326617.1	EST_HUMAN	hw05a11.x1 NCI_CGAP_Lu24 Homo saplens cDNA clone IMAGE:3182012 3'
2633	12500	22393	1.7	7.0E-31	7.0E-31 BE326517.1	EST HUMAN	hw05a11.x1 NCL_CGAP_Lu24 Homo saplens cDNA clone IMAGE:3182012 3'
9583	19168	25272	2.56	7.0E-31	X51755.1	TN	Human lambda-immunoglobulin constant region complex (germline)
3625	13539		2.79	8.0F-31	6 0F-31 AF223391 1	Ė	Homo saplens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial ods, alternatively spliced
6739	16618		3.68	6.0E-31	6.0E-31 AF055066.1	Į	Homo saplens MHC class 1 region
8120	18008	28255	2.11	6.0E-31	6.0E-31 AU119105.1	EST_HUMAN	AU119105 HEMBA1 Homo sapiens cDNA clorie HEMBA1005050 5
9191	18928	25351	1.88	6.0E-31	6.0E-31 AW372868.1	EST_HUMAN	RC6-BT0377-091299-031-D12 BT0377 Homo seplens cDNA
9319	19541		1.78	6.0E-31	6.0E-31 BE894488.1	EST_HUMAN	601433087F1 NIH_MGC_72 Hamo sapiens cDNA clone IMAGE:3918524 5'
185	10157	19973	2.85		5.0E-31 M60694.1	NT	Homo sapiens type I DNA topoisomerase genaļ exon 8
185	10157	19974	2.85		5.0E-31 M60694.1	LN.	Homo sepiens type I DNA topoisomerase gene; exon 8
!			,				7k06f04.x1 NCI_CGAP_GC8 Homo sapiens cDNA clone IMAGE:3443479 3' sImilar to TR:Q13537 Q13537
6915	16/93		1.2.1	5.0E-31		ESI HUMAN	SIMILAR TO FOCO ELEMENT: Contains LT.1 LT repetitive element;
581	10518		3.78	4.0E-31	4.0E-31 AJ271735.1	LN-	Homo saplens Xq pseudoautosomal region; segment 1/2
							POLYPEPTIDE N-ACETYLGALACTOSAMINYLTRANSFERASE (PROTEIN-UDP ACETYLGALACTOSAMINYLTRANSFERASE) (UDP-GALNAC:POLYPEPTIDE, N-
1594	11498	21357	1.04	4.0E-31	4.0E-31 Q10473	SWISSPROT	ACELYICALACTOSAMINYLTKANSTERASE (GALNACTT)
1776	11674		2.59	4.0E-31	4.0E-31 AL163280.2	LN.	Homo sapiens chromosome 21 segment HS21C080

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Probe SEQ ID NO:	Exen SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Vætue	Top Hit Acessian No.	Top Hit Detabase Source	Top Hit Descriptor
2759	12821		1.38	4.0E-31	5730038 NT	INT	Homo saplens SET domain and mariner transposase fusion gene (SETMAR) mRNA
9604	19186		1.52	4.0E-31	11430273 NT	INT	Homo sapiens KiAA0569 gene product (KIAA0569), mRNA
9718	19259		1.38	4.0E-31	4.0E-31 AB008681.1	TN	Homo saplens gene for activin receptor type IIB, complete cds
2550	12423	22314	1.42	3.0E-31	6005871 NT	LΝ	Homo sapiens SEC63, endoplasmic reticulum translocon component (S. cerevisiae) like (SEC63L), mRNA
88	16204	26366	9.03	3.0E-31	4826853 NT	L	Homo sapiens NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 8 (19kD, ASHI) (NDUFB8) mRNA
6425				3.0E-31		NT	Homo saplens hypothetical protein FLJ10842 (FLJ10842), mRNA
6745				3.0E-31	AL163206.2	N	Homo saplens chromosome 21 segment HS21C006
7525	17376	27585	2.86	3.0E-31	3.0E-31 D14523.1	۲	Horse mRNA for ferritin L-chain, complete cds
8028				3.0E-31	3.0E-31 P11174	SWISSPROT	40S RIBOSOMAL PROTEIN S15 (RIG PROTEIN)
8493	18366		96.36	3.0E-31	BF035327.1	EST_HUMAN	601458531F1 NIH_MGC_66 Home sapiens cDNA clone IMAGE:3862086 5
1875	١.	L	1.83	2.0E-31	AW838171.1	EST_HUMAN	QV2-LT0051-260300-111-f03 LT0051 Homo saplens cDNA
2167		21955	1.31	2.0E-31	AI393388.1	EST_HUMAN	tg44g05.x1 Soares_NFL_T_GBC_S1 Horno sapiens cDNA clone IMAGE:2111672.3'
2232	12174		2.18	2.0E-31	2.0E-31 AL119245.1	EST_HUMAN	DKFZp761G1513_r1 761 (synonym: hamy2) Homo sapiens cDNA clone DKFZp761G1513 5
7000	7,0000	70700		100	, , , , , ,	E C	aa88f11.s1 Stratagene fetal retina 937202 Homo saplens cDNA clone IMAGE:838413 3' similar to contains
I ASS	1		4.4	ZVE-31	ZUE-31 AA438824.1	ESI DOMAIN	ווואנג ווא ופלינות אינון
6502	16421	25483	99	20E-31 BE	BE350127.1	EST HUMAN	ht08g01.x1 NCI_CGAP_Kid13 Homo sapiens cDNA clone IMAGE:3146256 3' similar to contains MER29.b3 MER29 repetitive element ;
20.00			,	2000	4 4 0 7 7 7 0 4 4	TOT TOTAL	IND6F04.81 NCI_CGAP_C010 Homo sapiens cDNA clone IMAGE:1161055 3' similar to TR:Q13537 Q13537 MED27 TDANSDOSADI E EL EMENT COMPLETE CONCENSIS SECULENCE
227	L		ľ	2010		-1.	Leave in the control of the control
200	17550	2777	93.89	205-31	1001007 1001007 1001007 1001007	EST HIMAN	AV740048 Cti Homo sanjens CDNA clana Cti & Al R07 5
7700				20E-31	20E-31 AV710948.1	EST HUMAN	AV710948 Cu Homo sapiens cDNA clone CuAALB07 5
7677				20E-31	20E-31 BE408611.1	EST HUMAN	601304125F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3638310 5'
7677	17647		2.17	20E-31	20E-31 BE408611.1	EST_HUMAN	601304125F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3638310 5'
8292	18983		2	2.0E-31	AF148512.1	N	Homo sapiens hexoldnase II gene, promoter region
9431	19759		1.81	20E-31	20E-31 Al114527.1	EST_HUMAN	HA1110 Human fetal liver cDNA library Homo sapiens cDNA
							Homo saplens MAGE-B2 (MAGE-B2), MAGE-B3 (MAGE-B3), MAGE-B4 (MAGE-B4), and MAGE-B1
16			8.34	1.0E-31	1.0E-31 U93163.1	N	(MAGE-B1) genes, complete cds
1640	11544			1.0E-31	095371	SWISSPROT	OLFACTORY RECEPTOR 2C1
1640			2	1.0E-31	1.0E-31 095371	SWISSPROT	OLFACTORY RECEPTOR 2C1
1640				1.0E-31	095371	SWISSPROT	OLFACTORY RECEPTOR 2C1
4535	14428		1.01	1.0E-31	1.0E-31 AL134376.1	EST_HUMAN	DKFZp547B235_r1 547 (synonym: hfbr1) Homo saplens cDNA clone DKFZp547B235 5'

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					1	Т	Γ	12		9.13											0		ō, 4	to, 4					\neg
Top Hit Descriptor	DKFZp547B235_r1 547 (synonym: hfbr1) Homo sapiens cDNA clone DKFZp547B235 5'	MR3-ST0220-151299-028-a08_1 ST0220 Homo saplens cDNA	Homo sapiens minisatellite ceb1 repeat region	qf21h03.x1 NCI_CGAP_Brn25 Homo septens cDNA clone IMAGE:1750709 3' similar to TR:Q16595 O 46595 FRATAXIN	Human familial Alzheimer's disease (STM2) gene, complete cds	AV723976 HTB Homo sapiens cDNA clone HTBAAG01 5'	oz15a09.x1 Soares_fetal_liver_spleen_1NFLS_iS1 Homo sapiens cDNA clone IMAGE:1675384 3'	NUCLEAR ENVELOPE PORE MEMBRANE PROTEIN POM 121 (PORE MEMBRANE PROTEIN OF 121 KD) (P145)	Human chromosome 22 immunoglobulin V(K)I gene, part. with 5' breakpoint between orphon and neichbouring non-amplified region	tm34a10.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2159994 3' similar to contains MER29.13	MER29 repetitive element;	601511530F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3913087 5'	Homo sapiens PRO1181 mRNA, complete cds	Homo sapiens chromosome 21 segment HS21,C046	Homo saplens AT-binding transcription factor 1 (ATBF1), mRNA	Homo saplens AT-binding transcription factor 1 (ATBF1), mRNA	Homo sapiens FLH gene, partial	AV731500 HTF Homo sapiens cDNA clone HTFAKC07 5'	AV758634 BM Homo sapiens cDNA clone BMFBBH12 5'	AV758634 BM Homo sapiens cDNA clone BM/FBBH12 5'	2/95a07.s1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:448500 3' similar to contains THR.t3 THR repetitive element;	601156285F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3139701 5'	Homo sapiens myeloid/lymphoid or mixed-lineage leukemia (trithorax (Drosophila) homolog); translocated to, 4 (MLLT4) mRNA	Homo sapiens myeloid/lymphoid or mixed-lineage leukemia (trithorax (Drosophila) homolog); translocated to, 4	(MLLT4) mRNA	601156285F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3139701 5'	601173831F1 NIH_MGC_17 Homo saplens cDNA clone IMAGE:3529159 5'	H.sapiens mRNA for myosin	H.sapiens mRNA for myosin
Top Hit Database Source	EST_HUMAN	EST_HUMAN	NT	EST HIMAN	L	EST HUMAN	EST HUMAN	SWISSPROT	L		EST_HUMAN	EST_HUMAN	NT	LN	NT	NT	ΙN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	LN L		NT	EST_HUMAN	EST HUMAN	LN	NT
Top Hit Acession No.	AL134376.1	AW391679.1	AF048727.1	AINBRA34 1	150871.1	AV723976.1	AI056770.1	P52591	X17283 1		AI478104.1	BE888016.1	AF116627.1	AL163246.2	11432574 NT	11432574 NT	Y17293.1	AV731500.1	AV758634.1	AV758634.1	AA777621.1	BE279086.1	5174574 NT		5174574 NT	BE279086.1	BE296613.1		Z38133.1
Most Similar (Top) Hit BLAST E Value	1.0E-31	1.0E-31	1.0E-31		4-	_		7.05-32			6.0E-32	6.0E-32	5.0E-32	4.0E-32	4.0E-32	4.0E-32			3.0E-32	3.0E-32	3.0E-32								2.0E-32
Expression Signal	1.01	3.15	1.87	27.0	0 99	2.34	3.32	1.17	27.6		6.0	1.5	29.57	1.74	2.82	2.82	3.4	7.67	6.73	6.73	8.08	3.04	2.43		2.43	4.94	0.91	19.01	19.01
ORF SEQ ID NO:	24210	24927	25738	28411		26029		24426			22456	-	20779		26510	26511	20213	21210	27477	27478	28421		24882		24883		24454		25930
Exon SEQ (D NO:	14428	15159	15635	18187	12763	15905	11923	14639	18972		12568	16222	10936	10839	16342	16342	10392	11343	17271	17271	18176	18995	15090		15090	19314	14667		15806
Probe SEQ ID NO:	4535	5235	5728	8288	2835	800	2032	4754	8928		2702	6329	1018	915	8484	6484	84	1438	7404	7404	8297	8294	9655		9655	3802	4783	2800	2900

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Top Hit Descriptor	zn86c08.r1 Stratagene HeLa cell s3 937216 Homo sapiens cDNA clone IMAGE:563150 5'	zn86c08.r1 Stratagene HeLa cell s3 937216 Húmo sapiens cDNA clone IMAGE:563150 5'	AV736449 CB Hamo sapiens cDNA clone CBFBIA08 5	AV736449 CB Homo sepiens cDNA clone CBFBIA08 5	601573207F1 NIH_MGC_9 Homo saplens cDNA clone IMAGE:3834433 6'	Homo sapiens chromosome 11open reading frame 9 (C110RF9), mRNA	nw21g02.s1 NCI_CGAP_GCB0 Homo sepiens cDNA clone IMAGE:1241138 3' similar to contains THR.t3 THR repetitive element;	hw07c05.x1 NCI_CGAP_Lu24 Homo sapiens cDNA done IMAGE:3182216 3' similar to TR:O88539 O88539 WW DOMAIN BINDING PROTEIN 11.;	Homo sapiens calcium channel alpha1E suburit (CACNA1E) gene, exons 7-49, and partial cds, alternatively	spiiced	602021164F1 NC_CGAP_Bm67 Homo septens cDNA clone IMAGE:4156670 5	Homo sapiens chromosome 21 segment HS21C080	Homo sapiens short-chain elcohol dehydrogenisse family member (HEP27) mRNA	Homo saplens short-chain alcohol dehydrogenase family member (HEP27) mRNA	to12b09.x1 NCI_CGAP_Ut2 Homo sapiens cDNA clone IMAGE:2178809 3' similar to contains OFR.t1 OFR	NUTROLINGIAL	AV730056 H I F Homo sapiens cUNA cione H I FAVEUS 5	AV730015 HTF Homo sepiens cDNA clone H i FANF08 5	EST383398 MAGE resequences, MAGL Homo sapiens cDNA	602021164F1 NCI_CGAP_Brn67 Homo sepiens cDNA clone IMAGE:4156870 5'	EST383667 MAGE resequences, MAGL Homo eaplens cDNA	no16h01.s1 NCI_CGAP_Phe1 Homo saplensIcDNA clone IMAGE:1100881 3' similar to contains L1.t1 L1 repetitive element;	Homo sapiens chromosome 21 segment HS21 C085	Human glyceraldehyde-3-phosphate dehydrogenase (GAPDH) gene, complete cds	Homo sapiens similar to RAD23 (S. cerevisiae) homolog B (H. sapiens) (LOC63277), mRNA	Mus musculus SRY-box confaining gene 6 (Sox6), mRNA	Mus musculus SRY-box containing gene 6 (Sox6), mRNA	QV1-FT0189-100700-271-a02 FT0189 Homo saplens cDNA	Homo saplens solute carrier family 5 (choline fransporter), member 7 (SLC5A7), mRNA	Homo saplens spermidine synthase (SRM) mRNA	Homo sapiens spermidine synthase (SRM) mRNA		•
Top Hit Database Source	EST_HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	TN	EST HUMAN	EST HUMAN	!		EST_HUMAN	NT	TN	NT	100	LOUIS L	EST HOMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	Z	FZ	NT	NT	INT	EST_HUMAN	NT	NT	NT	İ	
Top Hit Acession No.	AA114294.1	AA114294.1	4736449.1	AV736449.1	BE743299.1	11439789	AA720574.1	BE327112.1		AF223391.1	BF347229.1	AL163280.2	5031736 NT	6031738 NT	180048	A1390113.1	AV730056.1	AV730015.1	AW971307.1	BF347229.1	AW971568.1	AA601416.1	AL163285.2	J04038.1	11429198 NT	6755609 NT	6755609 NT	BF373515.1	11141884 NT	4507208 NT	4507208 NT		
Most Similar (Top) Hit BLAST E Value	2.0E-32		_			1.0E-32	1.0E-32					9.0E-33	7.0E-33	7.0E-33	20 10	_		7.0E-33	7.0E-33	7.0E-33			-		6.0E-33	6.0E-33	6.0E-33	5.0E-33	5.0E-33	5.0E-33	5.0E-33		
Expression Signal	3.41	3.41	1.81	1.01	1.06	7.04	5.18	4.98		4.19	1.99	5.89	2.06	2.08		2.71	8.4	1.78	12.85	3.9	2.15	3.43	0.85	13.52	2.57	1.24	1.24	1.78	1.18	1.37	1.37		
ORF SEQ ID NO:	28884					26090	27065				27172		19850			70812		21439		L	28709			27059		27822	27823			21626			
Exan SEQ ID NO:	16694	L		19385		15958		13352		15773	16980	18065	1004	10041	1	12004		11573	l_,	18087	1	1	L	16865	l_	17600	17600	L	11735	L			
Probe SEQ ID NO:	6815	6815	8066	8908	3055	6198	6997	3435		5867	7103	8177	졌	2	17.7	CL17	2612	2800	3204	8203	8573	9274	3676	8869	7049	7750	7750	1739	1838	1856	1856		

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Table 4
Single Exon Probes Expressed in Heart

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	Top Hit Descriptor	Homo saplens chromosome 21 segment HS210085	Homo sapiens ubiquitin protein ligase E3A (human papilioma virus E6-associated protein, Angelman syndrome) (UBE3A), mRNA	Homo sapiens chromosome 21 segment HS21 C007	Homo sapiens RAB1, member RAS oncogene family (RAB1) mRNA	ab51b11.r1 Stratagene lung carcinoma 937218 Homo sapiens cDNA clone IMAGE:844317 5' similar to contains Atu repetitive element; contains MER28.b2 MER28 repetitive element;	Homo sapiens chromosome 21 segment HS21C010	UI-H-BI2-ahi-c-03-0-UI.s1 NCI_CGAP_Sub4 Homo saplens cDNA clone IMAGE:2727149 3	a71a08.r1 Stratagene colon (#937204) Homo sapiens cDNA clone IMAGE:510038 6' similar to gb:X12671_ma1 HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN A1 (HUMAN);	Homo sapiens killer cell immunoglobulin-like receptor, three domains, long cytoplasmic tail, 1 (KIR3DL1),	mRNA	ht08g01.x1 NCI_CGAP_Kid13 Homo saplens cDNA clone IMAGE:3146256 3' similar to contains MER29.b3 MER29 repetitive element :	ht09g01.x1 NCI_CGAP_Kid13 Homo saplens cDNA clone IMAGE:3146256 3' similar to contains MER29.b3	MER29 repetitive element;	AV647851 GLC Homo sapiens cDNA clone GLCBCF09 3'	qb67g03.x1 Soares_fetal_heart_NbHH19W Homo sepiens cDNA clone IMAGE:1705204 3' similar to contains OFR.t1 OFR repetitive element ;	MR0-HT0405-160300-202-d08 HT0405 Homo saplens cDNA	ab51g11.r1 Stratagene lung carcinoma 937218 Homo sapiens cDNA clone IMAGE:844388 5' sImilar to gb:X00734_cds1 TUBULIN BETA-5 CHAIN (HUMAN);	Homo sapiens hypothetical protein SIRP-b2 (SIRP-b2), mRNA	Homo sapiens hypothetical protein SIRP-b2 (SIRP-b2), mRNA	ql96d01.x1 Scares_NhHMPu_S1 Homo sapiens cDNA clone IMAGE:1880161 3'	ozz1d03.x1 Soares_fetal_liver_spleen_1NFL\$_S1 Homo saplens cDNA clone IMAGE:1675973 3' similar to gb:M29536 TRANSLATIONAL INITIATION FACTOR 2 BETA SUBUNIT (HUMAN);	Homo sapiens X-linked anhidroitic ectodermal dysplasia protein gene (EDA), exon 2 and flanking repeat	regions	QV3-BN0047-230200-102-b03 BN0047 Horrio sapiens cDNA	Human dystrophin (DMD) gene, exons 7, 8 and 9, and partial cds	wo88c06.x1 NCI_CGAP_Kid11 Homo sapieris cDNA clone IMAGE:2462410 3'
	Top Hit Database Source	LN		LN			N-	EST_HUMAN	EST_HUMAN			EST HUMAN		EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	LN	IN	EST_HUMAN	EST_HUMAN		뒫	EST_HUMAN	NT	EST_HUMAN
	Top Hit Acession No.	AL163285.2	11433063 NT	AL163207.2	4758987 NT	AA626621.1	١	AW 293349.1	AA053053.1		11425635 NT	BE350127.1		BE350127.1	AV647851.1	AI160189.1	BE159039.1	AA626883.1	11421332 NT	11421332 NT	AI277492.1	A1052256.1		AF003528.1	AW996818.1	U60822.1	Al927191.1
	Most Similar (Top) Hit BLAST E Value	5.0E-33	5.0E-33	4.0E-33/	-	4.0E-33		4.0E-33	4.0E-33		4.0E-33	3.05-33		3.0E-33	3.0E-33	2.0E-33	2.0E-33	2.0E-33		2.0E-33	2.0E-33	2.0E-33					1.0E-33
	Expression Signal	1.3	1.62	3.28	1.77	7.39	3.77	1.83	20.94		1.72	4 46		3.6	1.48	2.33	427	5.1	1.73	1.73	1.61	1.8		1.44	2.02	5.33	1.92
	ORF SEQ ID NO:	 -			21861		22265		25043		29102							24565		١						29024	
	Exan SEQ ID NO:	12111	l	1	i i		12373	14278	15239		18810	10980	1	10989	12721	10003	14219	14790	1	L	15775	17138		9994		18729	19151
	Probe SEQ ID NO:	2226	9082	1112	2078	2370	2498	4382	5318		2006	1073		1074	2400	66	4322	4911	5026	6026	5869	7281		80	8637	8921	9553

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	Top Hit Descriptor	Homo sapiens X-linked anhidroitic ectodermal dysplasia protein gene (EDA), exon 2 and flanking repeat regions	AV727809 HTC Hamo sapiens cDNA clane HTCCNC125	Homo sapiens Xq pseudoautosomal region; segment 1/2	yd15e05.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:108320 5'	y14c10.r1 Soares placenta Nb2HP Homo sapiens cDNA clone IMAGE:148722 5'	Human G2 protein mRNA, partial cds	Human G2 protein mRNA, partial cds	Mus musculus DAB/2J hair-specific (haci-1) gene	Homo sapiens Npw38-binding protein NpwBP (LOC51729), mRNA	Human splicing factor SRp55-1 (SRp-55) mRNA, complete cds	Rattus norvegicus putative four repeat Ion channel mRNA, complete cds	Homo sapiens mRNA for KIAA1435 protein, pertial cds	Homo sapiens chromosome 21 segment HS21,0009	tt94c06.x1 NCI_CGAP_Pr28 Homo sapiens cDNA clone IMAGE:2249194 3'	Homo sapiens hypothetical protein FLJ10989 (FLJ10989), mRNA	601468531F1 NIH_MGC_66 Homo sapiens cDNA clone IMAGE:3862086 5'	ADP,ATP CARRIER PROTEIN, LIVER ISOFØRM T2 (ADP/ATP TRANSLOCASE 3) (ADENINE NUCLEOTIDE TRANSLOCATOR 3) (ANT 3)	Homo sapiens X-linked anhidroitic ectodermal dysplasia protein gene (EDA), exon 2 and flanking repeat	regions	Homo sapiens WNT3 precursor (WNT3) mRNA, complete cds	Homo sapiens WNT3 precursor (WNT3) mRNA, complete cds	RC2-BT0506-240400-016-h08 BT0506 Homo sapiens cDNA	MR0-CT0068-280999-002-d11 CT0068 Hamo saplens cDNA	601484430F1 NIH_MGC_69 Homo sapiens cDNA clone IMAGE:3886999 6'	601484430F1 NIH_MGC_69 Homo sapiens cDNA clone IMAGE:3886999 5'	CM4-HT0193-061099-022-906 HT0193 Hamo sapiens cDNA	DKFZp564A1563_r1 564 (synonym: hfbr2) Homo sapiens cDNA clone DKFZp564A1563 5'	Homo saplens chromosome 21 segment HS21,0010	hh77b06.y1 NCI_CGAP_GU1 Hamo saplens cDNA clone IMAGE:2968787 5'	Homo saplens prohibitin (PHB) mRNA	naa33a08.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:3258134 3' similar to TR:075912 075912 DIACYLGLYCEROL KINASE IOTA. ;
, - -	Top Hit Database Source	FN	EST HUMAN	N	EST_HUMAN	EST_HUMAN	LN	ΙN	LΝ	۲	Ę	ΝΤ	NT.	ĮN.	EST_HUMAN	NT	EST_HUMAN	SWISSPROT		Ę	LΝ	TN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	LN	EST_HUMAN	6031190 NT	EST_HUMAN
	Top Hit Acession No.	AF003528.1	AV727809.1	AJ271735.1	T70845.1	H12866.1.	U10991.1	U10991.1	U03686.1	7706500 NT	U30883.1	AF078779.1	AB037856.1	AL163209.2	AI804667.1	8922807 NT	BF035327.1	P12236	į		AY009397.1	AY009397.1	BE071414.1	AW845706.1	BE874052.1	BE874052.1	AW368451.1	AL036635.1	AL163210.2	AW663302.1	6031190	F58993
	Most Similar (Top) Hit BLAST E Value	1.0E-33	1.0E-33	9.0E-34	7.0E-34	7.0E-34	6.0E-34	6.0E-34	6.0E-34	5.0E-34	5.0E-34		İ	5.0E-34	4.0E-34	4.0E-34	3.0E-34	1.0E-34		1.0E-34	1.0E-34	1.0E-34	1.0E-34	1.0E-34	1.0E-34	1.0E-34	1.0E-34	1.0E-34	1.0E-34	9.0E-35	8.0E-35	8.0E-35
	Expression Signal	3.04	1.34	2.09	1.98	3.08	1.46	1.46	1.4	2.23	4	1.18	2.12	2.01	2.11	0.98	4.81	8.56		1.48	0.78	0.78	4.02	0.86	1.99	1.89	3.87	8.89	2.92	1.26	24.25	3.3
	ORF SEQ ID NO:		25229		21198				25340		24631	27215			21735			.21254							25743	25744		27658		23286		21475
	SEQ (D NO:	9994	Ŀ	19399	11332	19020	10406	10406	18903	11734	14868	17021	17937	18446	11848	12552	18365	11393	L		13886	13886	14277	14789	15639	15639	1	17442	19307	13508	10189	
	Probe SEQ ID NO:	8723	9754	9927	1427	8340	463	463	9153	1837	4993	7144	8046	8578	1953	2687	8492	1488		3621	3979	3978	4381	4920	12/9	5731	7155	7591	9746	3594	219	1703

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				_	_												-	1880	<u>, ,, , , , , , , , , , , , , , , , , ,</u>	-7	(10 · Au		:	***		•	****	10 April 11
Top Hit Descriptor	nea33a08.x1 NCI_CGAP_Kid11 Homo saplens cDNA clone IMAGE:3289134 3' similar to TR:O75912 075912 DIACYLGLYCEROL KINASE IOTA.;	601809588F1 NIH_MGC_18 Homo sapiens cDNA clone IMAGE:4040324 5'	601236468F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3608513 5'	602184624T1 NIH_MGC_42 Homo sapiens cDNA clone IMAGE:4300660 3'	Homo saplens phosphatidylinositol glycan, class L (PIGL), mRNA	ah53h03.s1 Soares_testis_NHT Homo sepiens cDNA clone 1309397 3'	Homo saptens zinc finger protein 208 (ZNF208), mRNA	Homo saplens triple functional domain (PTPRF interacting) (TRIO), mRNA	Homo sapiens mRNA for KIAA1365 protein, partial cds	H.sapiens immunoglobulin kappa light chain variable region L14	Homo sapiens clk2 kinase (CLK2), propin1, cote1, glucocerebrosidase (GBA), and metaxin genes, complete	cas; metaxin pseudogene and glucocerebrosidase pseudogene, and thrombospondins (Tribos) gene, partall cds	601431984F1 NIH_MGC_72 Hamo saplens cDNA clone IMAGE:3917229 5	qg38c05.x1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1837448 3' similar to	SW:Y249_HUMAN Q92539 HYPOTHETICAL PROTEIN KIAA0249.;	qg38c05.x1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1837448 3' similar to SW:Y249 HUMAN Q92539 HYPOTHETICAL PROTEIN KIAA0249.	zh84f12.rf Soares_fetal liver_splean_1NFLS_S1 Homo sapiens cDNA clone IMAGE:428015 5	801109719F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3350405 5'	yu98a07.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:241236 5' similar to contains PTR5 recettive element :	601300705F1 NIH MGC 21 Homo sapiens cDNA clone IMAGE:3635401 5'	ht09g01.x1 NCI_CGAP_Kid13 Homo saplens cDNA clone IMAGE:3146256 3' similar to contains MER29.b3	MER29 repetitive element ;	DKFZp434L148_r1 434 (synonym: htes3) Homo saplens cDNA clane DKFZp434L148 5'	601125260F1 NIH_MGC_8 Homo sapiens cDNA clone IMAGE:3345063 5'	Homo sapiens phospholipid scramblase 1 gene, complete cds	MR1-TN0045-130900-010-e01 TN0045 Homo sapiens oDNA	7n25a09.xr NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3565361 3' similar to TR:Q9QZH7 Q9QZH7 F-BOX PROTEIN FBL2.;	7n25a09.x1 NOI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3565361 3' similar to TR:Q9QZH7 Q9QZH7 F-BOX PROTEIN FBL2.;
Top Hit Database Source	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	EST_HUMAN	NT	N	F	۲		Ļ	EST_HUMAN		EST_HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	FST HUMAN	EST HUMAN	I	EST_HUMAN	EST_HUMAN	EST_HUMAN	N	EST_HUMAN	EST_HUMAN	EST_HUMAN
Top Hit Acession No.	BF589937.1	8.0E-35 BF183195.1	BE378480.1	BF569282.1	11425417 NT	6.0E-35 AA757115.1	6005975 NT	6005921 NT	AB037786.1	5.0E-35 X63392.1		AF023268.1	5.0E-35 BE880992.1		A 208765.1	5.0E-35 Al208765.1	5.0E-35 AA001786.1	BE257907.1	H91193 1	4.0E-35 BE409102.1		BE350127.1	4.0E-35 AL046598.1	BE268182.1	AF224492.1	BF376402.1	BF433100.1	BF433100.1
Most Similar (Top) Hit BLAST E Value	8.0E-35 B	8.0E-35	8.0E-36	8.0E-35		6.0E-35	6.0E-35	6.0E-35				5.0E-35 A			5.0E-35 A	5.0E-35		İ	4 0F-35 F	4.0E-35				3.0E-35 B	3.0E-35	3.0E-35 Bi	3.0E-35 B	3.0E-35 B
Expression Signal	3.3	3.09	8	3.49	1.52	1.28	1.78	3.6	2.7	2.29		2.2	3.14		2.18	2.18	3.42	14.91	C.	1.29		1.82	6.84	9.63	2.42	1.06	22.73	22.73
ORF SEQ ID NO:	21478	24435	28217		25937	21154	21699	26876		21454		23992			26848	26849		21186	24552				27030			24465	24971	
Exan SEQ ID NO:	11604	14646	17966	18969	15811	11296	11820	16490	<u> </u>	11583		14208	L	L	16658	16658		L	11675		L.	16144	16835	11468	12167	14678	15197	15197
Probe SEQ ID NO:	1703	4761	8075	9265	5905	1391	1925	6610	7698	1681		4311	6761		6779	6779	8517	1415	1776	9042		6280	6957	1561	2283	4783	5275	5278

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Top Hit Descriptor	Homo sapiens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively spliced	K6932F Human fetal heart, Lambda ZAP Express Homo saplens cDNA clone K6932 5' similar to REPETITIVE ELEMENT	A971F Heart Homo sepiens cDNA clone A971,	Homo sapiens mRNA for Gab2, complete cds	Homo sapiens Grb2-associated binder 2 (KIAA0571), mRNA	Homo sapiens Grb2-associated binder 2 (KIAA0571), mRNA	Homo sapiens mRNA for KIAA0895 protein, partial cds	TCBAP2E4328 Pediatric pre-B cell acute lymphoblastic leukemia Baylor-HGSC project≂TCBA Homo sapiens cDNA clone TCBAP4328	TCBAP2E4328 Pediatric pre-B cell acute fumphoblastic leukemia Bayor-HGSC project=TCBA Homo sapiens	cDNA clone TCBAP4328	yq19a12.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:274079 5'	QV0-BT0701-210400-199-b04 BT0701 Homo saplens cDNA	H.saplens PROS-27 mRNA	Homo saplens Grb2-associated binder 2 (KIAA0571), mRNA	Homo sapiens Grb2-associated binder 2 (KIAA0571), mRNA	601496774F1 NIH_MGC_70 Homo saplens cDNA clone IMAGE:3898699 5'	601496774F1 NIH_MGC_70 Homo saplens cDNA clone IMAGE:3898699 5'	Homo sapiens chromosome 21 segment HS21C010	K6532F Human fetal heart, Lambda ZAP Express Homo saplens cDNA clone K6932 5' similar to REPETITIVE ELEMENT	fmfc16 Regional genomic DNA specific cDNA library Homo sapiens cDNA clone CR12-1	fmfc16 Regional genomic DNA specific cDNA library Homo saplens cDNA clone CR12-1	IL2-ST0162-131099-006-d12 ST0162 Homo sapiens cDNA	IL2-ST0162-131099-006-d12 ST0162 Homo sapiens cDNA	yd93a01.r1 Soares fetal liver spleen 1NFLS Homo saplens cDNA done IMAGE:115752 5' similar to SP:A44282 A44282 RETROVIRUS-RELATED POL POLYPROTEIN - HUMAN;	Homo sapiens hypothetical protein (LOC51233), mRNA	ht09g01.x1 NCI_CGAP_Kid13 Homo sapiens cDNA clone IMAGE.3146256 3' similar to contains MER29.b3 MER29 repetitive element :	ht09g01.x1 NC_CGAP_Kid13 Homo sapiens cDNA clone IMAGE:3146256 3' similar to contains MER29.b3 MER29 repetitive element :	
Top Hit Database Source	TN	EST_HUMAN	EST_HUMAN	NT	TN	LN.	LΝ	EST HIMAN	NUMBER OF THE PROPERTY OF THE	EST_HUMAN			NT	TN		EST_HUMAN	EST_HUMAN	TN	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	N-	EST_HUMAN	EST_HUMAN	
Top Hit Acession No.	AF223391.1	V88965.1	T11909.1	AB018413.1	6912459 NT	6912459 NT	AB020702.1	3E247575 4		BE247575.1	H49239.1	BF332417.1	X59417.1	6912459 NT	6912459 NT	3E904978.1	BE904978.1	AL163210.2	N88965.1	AA631949.1	AA631949.1	AW389473.1	AW389473.1	187947.1	7705994 NT	BE350127.1	BE350127.1	
Most Similar (Top) Hit BLAST E Value	3.0E-35	2.0E-35	2.0E-35	2.0E-35	2.0E-35	2.0E-35	2.0E-35	2 AE.3A	2:01-00	2.0E-35	2.0E-35	2.0E-35	2.0E-35	2.0E-36	2.0E-35	2.0E-35	2.0E-35	2.0E-35	2.0E-35	1.0E-35	1.0E-35	1.0E-35	1.0E-35	1.0E-35		1.0E-35	1.0E-35	
Expression Signal	1.81	1.74	1.25	5.2	76.0	76.0	0.88	•	60:	1.09	2.55	1.66	3.72	1.38	1.36	1.51	1.61	3.97	4.17	4.38	4.38	44.43	44.43	1.18	2.31	1.11	1,11	
ORF SEQ ID NO:		19902	20928			22992		22530	20002	23540		25401	28312	22891	22892	25354	25355		19902		19829	20500	20501		22262	22498		
Exan SEQ ID NO:	17325	12659	11083	12058	13193	13193	13427	197.47		13747	14462	15347	18083	13183	13193	18936	18936	19264	12659	10028	10028	10667	10667	10817	L	12602		
Probe SEQ ID NO:	7465	183	1171	2171	3272	3272	3511	2695	3	3835	4570	5428	8175	8028	8028	8205	9208	9725	9832	\$	8	735	735	891	2495	2740	2740	

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Top Hit Descriptor	Homo saplens transcription elongation factor B (SIII), polypeptide 1-like (TCEB1L) mRNA	AV650422 GLC Hamo sapiens cDNA clone GLCCEF06 3'	AV650422 GLC Hamo sapiens cDNA clone GLCCEF06 3'	Mus musculus activin receptor interacting protein 1 (Arip1-pending), mRNA	Mus musculus activin receptor interacting protein 1 (Arip1-pending), mRNA	Homo sapiens chromatin assembly factor 1, subunit B (p60) (CHAF1B), mRNA	AU158595 PLACE3 Hamo sapiens cDNA clone PLACE3000382 3'	AU158595 PLACE3 Homo saplens cDNA clone PLACE3000382 3'	promma-7.D01.r bytumor Homo saplens cDNA(5'	Homo saplens hypothetical protein (LOC51233), mRNA	Homo saplens casein kinase 1, epsilon (CSNK1E), mRNA	601584833F1 NIH_MGC_7 Homo sapiens aDINA clane IMAGE:3938985 5'	CM1-CT0315-091289-063-d07 CT0315 Homo saplens cDNA	Homo sapiens C-terminal binding protein 2 (CTBP2) mRNA	Human carcinoembryonic antigen gene family member 12 (CGM12) gene, exons L and L/N	Human carcinoembryonic antigen gene family member 12 (CGM12) gene, exons L and L/N	Homo saplens glutathione transferase A4 gene, exon 1	Homo saplens ninjurin 2 (NINJ2), mRNA	Homo sapiens TCL8 gene, exon 12	UI-H-BW1-anv-c-12-0-UI.s1 NCI_CGAP_Sub7 Homo sepiens cDNA clone IMAGE:3083542 3'	thg3b06.x1 Soars_NSF_F8_9W_OT_PA_P_51 Homo saplens cDNA clone IMAGE:2126195 3' similar to gb:M1949 PANCREATIC SECRETORY TRYPSIN INHIBITOR PRECURSOR (HUMAN);	hoo6h02.x1 NCI_CGAP_Co14 Homo sepiens cDNA clone IMAGE:3036927 3' similar to SW:IMA2_HUMAN	PSZZBZ IMPORTIN ALTRA-Z SUBONTI,	Homo sapiens syncylin precursor, mikiva, complete cos	#85c09.x1 NCI_CGAP_CLL1 Home sepiens cDNA clone IMAGE:2107024 3' similar to contains MER9.b2	MER9 repetitive element;	601305064F1 NIH_MGC_39 Homo sapiens cDNA clane IMAGE:3639782 5'	Homo saplens Xq pseudoautosomal region; segment 1/2	601285567F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3607289 5'	Homo saplens chromosome 21 segment HS21,0009	Homo sapiens API5-like 1 (API5L1), mRNA	Homo sapiens API5-like 1 (API5L1), mRNA	Homo saplens Xq pseudoautosomal region; segment 1/2
Top Hit Database Source	NT	EST_HUMAN	EST_HUMAN	TN	NT	IN	EST_HUMAN	EST_HUMAN	EST_HUMAN	TN	NT	EST_HUMAN	EST_HUMAN	TN	IN	N	NT	TN	N	EST_HUMAN	EST HUMAN		EST HUMAN	Z		EST_HUMAN	EST_HUMAN	IN	EST_HUMAN	ΙN	۲	L	INT
Top Hit Acession No.	6006030 NT	4V650422.1	4V650422.1	7656905 NT	7656905 NT	11526236 NT	AU158595.1	AU158595.1	41625119.1	7705994 NT	11418110 NT	3E792832.1	4W857579.1	4557498 NT	J06672.1	J06672.1	AF052051.1	7706622 NT	4B035346.1	3F515101.1	A1435169.1		AW 780143.1	AF208161.1		AI380499.1	BE737154.1	AJ271735.1	3E388436.1	AL163209.2	6729729 NT	5729729 NT	4,1271735.1
Most Similar (Top) Hit BLAST E Value	1.0E-35	1.0E-35	1.0E-35	1.0E-35	1.0E-35	1.0E-35	1.0E-35	1.0E-35	1.0E-35	1.0E-35	1.0E-35	1.0E-35	7.0E-36	7.0E-36	7.0E-36	7.0E-36	7.0E-36	6.0E-36	8.0E-36	6.0E-38	6.0E-36		6.0E-36	6.0E-36		6.0E-36	6.0E-36	6.0E-36	5.0E-38	5.0E-36	5.0E-38	5.0E-36	5.0E-36
Expression Signal	1.07	2.2	2.2	4.7	4.7	1.43	2	2	4.16	1.69	1.37	2.28	0.94	4.03	6.04	6.04	5.23	1.88	6.17	0.91	19.79		3.34	2.21		2.74	1.69	69.9	1.71	1.96	1.42	1.42	2.64
ORF SEQ ID NO:	22826	22848		24003	24004	26154	27578	27679		22262			22622		26548	26549	25279		-	23280				27098		28931	25071	19928	22478	23264		24359	Ц
Exon SEQ ID NO:	13030	13051	13051	14222	14222	15301	19469	19469	18782	12369	18971	19198	12827	13007	16371	16371	19071	11854	12247	13501	(1	16087	16904		18646	19580	10107	12584	13475	14564	1	Li
Probe SEQ ID NO:	3104	3126	3128	4325	4325	5382	7501	7501	8977	9186	9267	9620	2800	3080	6512	6512	9423	1959	2367	3587	5268		6221	7027		8833	9821	133	2722	3561	4678	4678	8026

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wb37c12x1 NCI_CGAP_GC6 Homo sapiens cDNA clone IMAGE:2307862 3' similar to contains Alu Homo saplens a disintegrin and metalloproteinase domain 22 (ADAM22), trancript variant 3, mRNA RETROVIRUS-RELATED POL POLYPROTE:IN [CONTAINS: REVERSE TRANSCRIPTASE 2051a12.r1 Stratagene endothelial cell 937223 Homo saplens cDNA clone IMAGE:590398 5' 2051a12.r1 Stratagene endothelial cell 937223 Homo saplens cDNA clone IMAGE:590398 5' yc44a07.11 Stratagene liver (#937224) Homo sapiens cDNA clone IMAGE:83508 5' 601300938F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3635480 5' Homo sapiens nuclear factor of activated T-cells, cytoplasmic 2 (NFATC2), mRNA AV753629 TP Homo sapiens cDNA clone TPGABH01 5 EST06648 Infant Brain, Bento Soares Homo sapiens cDNA clone HIBBJ28 5' end xe82b07.x1 NCI_CGAP_Brn35 Homo sapiens cDNA clone IMAGE:2614357 31 QV3-NN1023-010600-199-h01 NN1023 Homo sapiens cDNA zu69c10.r1 Soares_testis_NHT Homo sepiens cDNA clone IMAGE:743250 5 2820020.5prime NIH_MGC_7 Homo sapiens CDNA clone IMAGE.2820020 5 601298574F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:3628386 5 601282266F1 NIH_MGC_44 Homo sapiens cONA clone IMAGE:3604168 5' 601282266F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3604168 5 601458531F1 NIH_MGC_66 Hamo sapiens cDNA clone IMAGE:3862086 67 601106343F1 NIH_MGC_16 Hamo sapiens cDNA clone IMAGE:3342706 67 602136493F1 NIH_MGC_83 Homo sapiens cDNA clone IMAGE:4272886 5 Homo sapiens DNA for amyloid precursor protein, complete cds Homo sapiens DNA for amyloid precursor protein, complete cds Top Hit Descriptor Homo sapiens calcineurin binding protein 1 (KIAA0330), mRNA PM3-BN0176-100400-001-g04 BN0176 Homo sapiens cDNA QV0-OT0030-240300-174-h04 OT0030 Homo sepiens cDNA RC1-HT0217-131199-021-h07 HT0217 Homo sapiens cDNA RC1-HT0217-131199-021-h07 HT0217 Homo sapiens cDNA Human platelet Glycoprotein IIb (GPIIb) gene, exons 2-29 Homo sapiens KIAA0952 protein (KIAA0952), mRNA Mus musculus junctophilin 1 (Jp1-pending), mRNA Homo saplens neurexin III-alpha gene, partial ods Mus musculus p47-phox gene, complete cds ENDONUCLEASE repetitive element; EST_HUMAN EST_HUMAN HUMAN EST HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN **EST HUMAN** EST HUMAN EST HUMAN EST_HUMAN EST_HUMAN EST_HUMAN **EST HUMAN** EST_HUMAN EST_HUMAN Top Hit Database HUMAN SWISSPROT **EST HUMAN** EST_HUMAN Source EST 눋 눋 11417862 10181139 Top Hit Acession 4.0E-36 AV753629.1 BE382574.1 BE010038.1 AW247772.1 BE389299.1 4.0E-36 BE389299.1 4.0E-36 AA400370.1 AF099810.1 BE409310.1 BF673761.1 .0E-36 AA148034.1 1.0E-36 AA148034.1 1.0E-38 AW103658.1 BE146523.1 1.0E-36 BF364169.1 ģ 1.0E-36 AI867714.1 4.0E-36 M33320.1 D87675.1 T08756. 4.0E-36 2.0E-38 4.0E-36 4.0E-36 4.0E-36 4.0E-36 3.0E-36 1.0E-38 4.0E-36 .0E-36 2.0E-36 1.0E-36 6.0E-36 3.0E-36 2.0E-36 (Top) Hit BLAST E Aost Similar Value 8 8 8 1.58 2.19 2.85 0.89 5.15 2.16 0.86 5.94 2.36 3.21 3.21 1.74 3.99 0.86 4. Expression Signal 26711 ORF SEQ ID NO: 21196 23036 23037 25704 26547 27050 27051 28584 24534 20643 21884 27287 25337 20961 21389 28481 24081 22857 25127 19545 15660 16520 19010 15483 16620 11530 13231 16370 18233 12136 14297 18325 10793 SEQ ID 13231 15602 16856 16856 10614 15291 11987 13057 11987 12043 Ĕ ÿ 5752 Probe SEQ ID 9318 1424 3310 3310 5693 6979 6979 8356 9334 9380 2252 4402 8452 3132 5567 5962 2098 2155 8840 6640 7220 1205 1626 6511 6371 4877 867 681 ÿ

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Top Hit Descriptor	CM3-NN0081-140400-147-h12 NN0081 Homo sapiens cDNA	UI-HF-BN0-ale-c-03-0-UI.11 NIH_MGC_50 Homo saplens cDNA done IMAGE:3079277 5'	Homo saplens Ran GTPase activating protein 1 (RANGAP1), mRNA	Homo sapiens chromosome 22 open reading frame 2 (C22ORF2), mRNA	Homo sapiens chromosome 21 segment HS21C013	Homo saplens Sad1 unc-84 domain protein 2 (SUN2) mRNA, partial cds	ws80b07.x1 NCI_CGAP_Co3 Homo sapiens cJNA clone IMAGE:2504245 3'	ws80b07.x1 NCI_CGAP_Co3 Homo sapiens cDNA clone IMAGE:2504245 3'	73D4 Human retina cDNA Tsp509I-cleaved subilbrary Homo saplens cDNA not directional	Homo sapiens mRNA for KIAA0877 protein, partial cds	CM0-UT0003-050800-503-d09 UT0003 Homo sapiens cDNA	ht09g01.x1 NCI_CGAP_Kid13 Homo sapiens cDNA clone IMAGE:3146256 3' similar to contains MER29.b3 MER29 repetitive element;	ht09g01.x1 NCI_CGAP_Kid13 Homo sapiens cDNA clone IMAGE:3146256 3' similar to contains MER29.b3	MERZ9 repetitive element;	RC1-CN0008-210100-012-a09_1 CN0008 Homo sapiens cDNA	H.sapiens DMA, DMB, HLA-Z1, IPP2, LMP2, TAP1, LMP7, TAP2, DOB, DQB2 and RING8, 9, 13 and 14 denes	DKET 1944 61 194 (evanorum: http:// Homo canions a DNA alana DKET 1949 450420 6.	wk25b11 x1 NCI CGAP Brn25 Home sapiens cDNA clone IMAGE:2413341 3' similar to contains PTR5 t2	PTR5 repetitive element;	tm87g03.x1 NCI_CGAP_Bm25 Homo sapiens cDNA clone IMAGE:2165140 3' similar to contains L1.b3 L1	repetitive element;	Homo sapiens Sad1 unc-84 domain protein 2 (SUN2) mRNA, partial cds	EST178035 Colon carcinoma (HCC) cell line Homo saplens cDNA 5 end	EST178035 Colon carcinoma (HCC) cell line Homo sapiens cDNA 5' end	Homo sapiens glycine C-acetyltransferase (2-amino-3-ketobutyrate-CoA ligase) (GCAT), mRNA	Homo sapiens NOD1 protein (NOD1) gene, exons 1, 2, and 3	zi90b04.s1 Soares_fetal_liver_spleen_1NFLS_iS1 Homo sapiens cDNA clone IMAGE:4480153'	EST62910 WATM1 Homo sapiens cDNA clone 62910 similar to human STS G04101	DKFZp434L2418_r1 434 (synonym: htes3) Homo saplens cDNA clone DKFZp434L2418	DKFZp434L2418_r1 434 (synonym: htes3) Hojno sapiens cDNA clone DKFZp434L2418	EST373222 MAGE resequences, MAGF Homb sapiens cDNA	EST373222 MAGE resequences, MAGF Homb saplens cDNA
Top Hit Database Source	EST_HUMAN	EST_HUMAN	NT	FZ	N	LΝ	EST_HUMAN	EST_HUMAN	EST_HUMAN	TN	EST_HUMAN	EST HUMAN		EST_HUMAN	EST_HUMAN	LΝ	LI INVAN	NUMBER	EST_HUMAN		T_HUMAN	NT	EST_HUMAN	EST_HUMAN	NT	TN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN		EST_HUMAN
Top Hit Acession No.	AW897636.1	AW 504143.1	11418177	11418121 NT	AL163213.2	AF202723.1	AW009277.1	AW009277.1	W22618.1	AB020684.1	BE698077.1	BE350127.1		BE350127.1	AW840840.1	X87344.1	1 000000	1.04200.1	AI817700.1		AI536702.1	AF202723.1	AA307123.1	AA307123.1	7657117 NT	AF149773.1	AA702794.1	N62051.1	AL048956.1	AL048956.1	AW961150.1	AW961150.1
Most Similar (Top) Hit BLAST E Value	1.0E-36		1.0E-36	1.0E-36	1.0E-36	1.0E-36		9.0E-37	9.0E-37	8.0E-37	8.0E-37	8.0E-37	-	_	8.0E-37	8.0E-37		-	7.0E-37		7.0E-37	6.0E-37	5.0E-37	6.0E-37	5.0E-37		4.0E-37	4.0E-37	3.0E-37			3.0E-37
Expression Signal	3.43	3.91	3.74	1.29	3.07	2.89	1.96	1.96	2.79	1.38	1.7	4.1		4.1	5.63	6.25	2 64	2	6.77		4.16	2.48	3.37	3.37	4.17	3.57	2.23	1.11	2.85	2.85	1.7	3.02
ORF SEQ ID NO:	28444	28857		25305			26390	26391		24781		25547		25548	25576	28670			28274		28388		25716	25717			22145		21755	21756		
Exon SEQ ID NO:	18194	18574	18934	19040	19214	18371		16231	19093	15010	15136	15475	l	J	15499	16482	11170		18028		18148	19292	15615	15615	18171			15027		11863	L	12863
Probe SEQ ID NO:	8317	8686	9203	9374	9647	0686	සෙසෙ	6368	9462	5143	6213	5559		5558	5584	6802	1282	3	81.40		8268	9775	5707	2029	8282	9198	2374	5160	1970	1970	2465	2936

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Table 4
Single Exon Probes Expressed in Heart

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Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
10980		1.94	2.0E-37	AU131202.1	EST_HUMAN	AU131202 NT2RP3 Homo saplens cDNA clone NT2RP3002166 5'
10980		1.94	2.0E-37	AU131202.1	EST_HUMAN	AU131202 NT2RP3 Homo sapiens cDNA clone NT2RP3002166 5
11816	21695	1.47	2.0E-37	\L163247.2	INT	Homo sapiens chromosome 21 segment HS21C047
13730	23519	5.05	2.0E-37	4503210 NT	NT	Homo sapiens cytochrome P450, subfamily XXVIIA (steroid 27-hydroxylase, cerebrotendinous xanthomatosis), polypeptide 1 (CYP27A1b) mRNA
15912	26039	3.36	2.0E-37	2.0E-37 AA346720.1	EST_HUMAN	EST52831 Fetal heart II Homo sepiens cDNA 5' end
16565	28759	3.23	2.0E-37 B	F204032.1	EST_HUMAN	601869157F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:4111406 5'
18657	28945	18.4	2.0E-37 A	AF176013.1	NT	Homo sapiens J domain containing protein 1 isororm b (JDP1) mRNA, complete cds
19403		3.15		11417972	LN	Homo sapiens pescadilio (zebrafish) homolog 1, containing BRCT domain (PES1), mRNA
11932	21828	3.61	1.0E-37	AL163281.2	TN	Homo sapiens chromosome 21 segment HS21C081
13789	23577	22.51	1.0E-37	AF189011.1	NT	Homo sapiens ribonuclease III (RN3) mRNA, complete cds
13974	23753	96.0	1.0E-37	BE872365.1	EST_HUMAN	601448619F1 NIH_MGC_65 Homo sapiens cDNA clone IMAGE:3852652 5'
14737		2.13	1.0E-37 B	F371719.1	Г	QV0-FN0180-280700-318-c10 FN0180 Homo sapiens cDNA
						zp21b02.r1 Stratagene neuroepithelium (#937231) Homo saplens cDNA clone IMAGE:610059 5' similar to
16949		2.85	1.0E-37	1.0E-37 AA171406.1	EST_HUMAN	contains L1.t2 L1 repetitive element ;
17973	28222	20.59	1.0E-37	122878.1		Human somatic cytochrome c (HC1) processed pseudogene, complete cds
19122		2.46	1.0E-37 B	E771814.1	EST_HUMAN	CM3-FT0096-140700-243-d07 FT0096 Homo sapiens cDNA
18447		3.05	9.0E-38	10048482 NT	NT	Rattus norvegicus multidomain presynaptic cytomatrix protein Piccolo (LOC56768), mRNA
11113		1.95	8.0E-38	11436955 NT		Homo saplens Grb2-associated binder 2 (KIAA0571), mRNA
12326			8.0E-38	BF346221.1	T_HUMAN	602018401F1 NCI_CGAP_Brn87 Homo sepiens cDNA clone IMAGE:4153992 5
11113		1.36	8.0E-38	11436955		Homo sapiens Grb2-associated binder 2 (KIAA0571), mRNA
12023	21820	1.38	7.0E-38	7.0E-38 AW972825.1	EST_HUMAN	EST384920 MAGE resequences, MAGL Homo saplens cDNA
12933	22726	1.78	6.0E-38 B	F033033.1	EST_HUMAN	601455722F1 NIH_MGC_68 Homo sapiens cDNA clone IMAGE:3859348 5'
15352	25407	1.69	6.0E-38	11425114 NT		Homo sapiens zinc finger protein ZNF287 (ZNF287), mRNA
15352	25408	1.69	6.0E-38	11425114 NT	N⊤	Homo sapiens zinc finger protein ZNF287 (ZNF287), mRNA
18842		4.48	6.0E-38	11435947 NT	NT	Homo sapiens chromosome 12 open reading frame 3 (C12ORF3), mRNA
19143	25288	99.9	8.0E-38	AB002059.1	TN	Homo sapiens DNA for Human P2XM, complete cds
10642	20468	1.15	5.0E-38	W971819.1	EST_HUMAN	EST383908 MAGE resequences, MAGL Homo saplens cDNA
12281	22178	1.79	6.0E-38	6.0E-38 AJ237740.1	NT	Homo sapiens RiBilR gene (partial), exon 8
16076	26225	2.42	5.0E-38	5.0E-38 BE871610.1	T_HUMAN	601450148F1 NIH_MGC_65 Homo sapiens cDNA clone IMAGE:3854074 5'
10092	18309	2.56	4.0E-38	4.0E-38 Z25466.1	LN	B.taurus mitochondriai aspartate aminotransferase mRNA, complete CDS
10092	19910	2.58	4.0E-38	4.0E-38 Z25466.1	NT	B.taurus mitochondrial aspartate aminotransferase mRNA, complete CDS
11055	20897	0.82	3.0E-38	11435947		Homo saplens chromosome 12 open reading frame 3 (C12ORF3), mRNA
11943		4.39	3.0E-38	F003530.1	IN	Homo sapiens homeobox protein CDX4 (CDX4) gene, complete cds and flanking repeat regions

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	Top Hit Descriptor	Homo sapiens HIRA interacting protein 4 (dnalike) (HIRIP4), mRNA	SSU72 PROTEIN	SSU72 PROTEIN	601157633F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3504272 5'	Homo saplens chromosome 21 segment HS21C100	CM3-FT0181-140700-241-f07 FT0181 Homo sapiens cDNA	yw88b04.r1 Soares melanocyte 2NbHM Homo capiens cDNA clone IMAGE:249775 5	w88b04.r1 Soares melanocyte 2NbHM Homo sapiens cDNA clone IMAGE:249775 5'	Homo saplens chromosome 21 segment HS21C048	Homo sapiens chromosome 21 segment HS21C048	Homo sapiens SMT3 (suppressor of mif two 3, yeast) homolog 2 (SMT3H2), mRNA	zw30d01.r1 Soares ovary tumor NbHOT Homo saplens cDNA clone IMAGE:770785 5' similar to SW:MA12_RABIT P45701 MANNOSYL-OLIGOSACCHARIDE ALPHA-1,2-MANNOSIDASE;	zw30d01.r1 Soares ovary tumor NbHOT Homo sepiens cDNA clone IMAGE:770785 5' similar to SW:MA12 RABIT P45701 MANNOSYL-OLIGOSACCHARIDE ALPHA-1,2-MANNOSIDASE :	MR3-HT0487-150200-113-g01 HT0487 Homo sapiens cDNA	hu09g02.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3186130 3' similar to TR:002710 002710 GAG POLYPROTEIN;	Homo saplens mRNA for KIAA0145 protein, partial cds	QV2-HT0698-080800-293-a05 HT0698 Homo saplens cDNA	Homo saplens leucine-rich repeat-containing G protein-coupled receptor 6 (LGR6) mRNA, partial cds	Homo sapiens leucine-rich repeat-containing G protein-coupled receptor 6 (LGR6) mRNA, partial cds	AV726988 HTC Homo sapiens cDNA clone HTCAXH07 5'	Homo saplens gene for kinesin-like protein, complete cds	CHR220580 Chromosome 22 exon Homo sepiens aDNA alone C22_788 6'	E1 beta=pyruvate dehydrogenase beta (promoter) [human, placenta, Genomic, 1280 nt]	Homo sapiens sulfotransferase-related protein (SULTX3), mRNA	zu62b02.r1 Soares_testis_NHT Homo sapiens oDNA clone IMAGE:742639 6' similar to contains element MER19 repetitive element;	Homo sapiens guanine nucleotide binding protein-like 1 (GNL1), mRNA	Homo sapiens KIAA0173 gene product (KIAA¢173), mRNA	Homo sapiens cyclin K (CCNK) gene, expn 7
	Top Hit Database Source	N _T	SWISSPROT	SWISSPROT	EST_HUMAN	IN	EST_HUMAN	EST_HUMAN	EST_HUMAN	LN LN	NT	TN	EST_HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	NT	EST_HUMAN	LΝ	FN	EST_HUMAN	NT	EST_HUMAN	NT	NT	EST HUMAN	L	TN	NT
	Top Hit Acession No.	7549807 NT	P53538	P53538	BE279301.1	AL163300.2	BF373664.1	H85494.1	H85494.1	AL163248.2	AL163248.2	5902097 NT	AA437353.1	AA437353.1	BE165980.1			BE712790.1	AF190501.1	AF190501.1	AV726988.1	AB012723.1	H55641.1	S74906.1	11418248 NT	AA401570.1	4885288 NT	7681969 NT	AF270831.1
	Most Similar (Top) Hit BLAST E Value	3.0E-38	3.0E-38	3.0E-38	3.0E-38	3.0E-38	3.0E-38	3.0E-38	3.0E-38	3.05-38	2.0E-38	2.0E-38	2.0E-38	2.0E-38	-	2.0E-38	2.0E-38		2.0E-38	2.0E-38	2.0E-38	2.0E-38	2.0E-38		2.0E-38	1.0E-38	1.0E-38	1.0E-38	1.0E-38
	Expression Signal	1.11	1.85	1.65	0.85	7.17	7.64	1.78	1.78	1.58	1.41	2.6	1.68	1.88	4.7	1.47	1.66	5.24	3.69	3.69	4.05	2	3.86	2.28	2.58	2.29	0.94	Ŧ	1.58
	ORF SEQ ID NO:		23480	23481					27095		19838	21120	21390	21391			28063	28885	28008	29009			25313				21736		22221
	Exan SEQ ID NO:	13552	13693	13693	14406			16902	16902	17577	10032	11264	11531	11531	16819	17182	17820	18596	18715	18715	18876	18878	19064	19099	19336	10993	11849		12322
	Probe SEQ ID NO:	3638	3781	3781	4513	6050	6478	7025	7025	77.27	44	1358	1627	1627	. 6941	7316	7970	8781	8907	8907	9112	9115	9412	9472	9824	1077	1954	1973	2445

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. Top Hit Descriptor	Homo sapiens mRNA for KIAA1442 protein, partial cds	Homo saplens low density lipoprotein receptor-jelated protein 6 (LRP6) mRNA, and translated products	Homo saplens chromosome 21 segment HS21,0003	Homo sapiens chromosome 21 segment HS21,0003	Homo saplens hypothetical protein FLJ10600 (FLJ10600), mRNA	Mus musculus otogelin (Otog), mRNA	Mus musculus otogelin (Otog), mRNA	Homo sapiens mRNA for KIAA0612 protein, peirtial cds	ht09g01.x1 NCI_CGAP_Kid13 Homo saplens :DNA clone IMAGE:3146256 3' similar to contains MER29.b3 MER29 repetitive element ;	Homo sapiens chromosome 21 segment HS21,0084	Homo sapiens ATPase, H+ transporting, lysosomal (vacuolar proton pump) 16kD (ATP6C) mRNA	Homo sapiens estrogen receptor-binding fragment-associated gene 9 (EBAG9) mRNA	wh53f10.x1 NCI_CGAP_Kld11 Homo saplens cDNA clone IMAGE:2384491 3' similar to TR:P87890 P87890 PDD PDD PDD CEIN :	Homo sapiens chromosome 21 segment HS21C027	QV1-BT0631-040900-357-f02 BT0631 Homo sapiens cDNA	7634c03.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3284356 3' similar to WP:R151.6	CE00828;	Homo sapiens X-linked anhidroitic ectodermal dysplasia protein gene (EDA), exon 2 and flanking repeat regions	a(36b04.x1 Barstead coton HPLRB7 Homo sepiens cDNA clone IMAGE:2374063 3' similar to TR:Q15408	Q15408 NEUTRAL PROTEASE LARGE SUBUNIT; contains LTR7.tt LTR7 repetitive element;	Homo sapiens hypothetical protein FLJ10803 (FLJ10803), mRNA	Chlorocebus aethiops mRNA for ribosomal protein S4X, complete cds	Homo saplens chromosome 21 segment HS21 C010	ae92g04.s1 Stratagene schizo brain S11 Horns sapiens cDNA clone IMAGE:1020438 3' similar to contains	OFR.b1 OFR repetitive element;	Homo saplens Ran GTPase activating protein (RANGAP1), mRNA	QV0-FN0063-260600-278-c06 FN0063 Hamo sapiens cDNA	fmfc16 Regional genomic DNA specific cDNA/library Homo sapiens cDNA clone CR12-1	firifc16 Regional genomic DNA specific cDNA library Homo sapkans cDNA clone CR12-1
Top Hit Database Source	NT	TN	NT	FZ	NT	NT L	LΝ	ΝΤ	EST_HUMAN	NT	NT.	N	MAAN ILI	TN TN	EST HUMAN		EST_HUMAN	ħ		EST_HUMAN	N	FN	N		EST_HUMAN	ΓN	EST_HUMAN	EST_HUMAN	EST_HUMAN
Top Hit Acession No.	AB037863.1	4505016 NT	AL163203.2	AL163203.2	8922543 NT	7305360 NT	7305360 NT	4B014512.1	3E350127.1	AL163284.2	4502312 NT	4758229 NT	100000	41 163227 2	3F331829 1		3E670394.1	AF003528.1		AI750154.1	11420289 NT	AB015610.1	AL163210.2		AA682949.1	11418177 NT	BE836452.1	AA631949.1	AA631949.1
Most Similar (Top) Hit BLAST E Value	1.0E-38	1.0E-38	1.0E-38	1.0E-38	1.0E-38	1.0E-38	1.0E-38	1.0E-38	1.0E-38	1.0E-38	8.0E-39	8.0E-39	100	7.0F-39	6.0F-39		6.0E-39	6.0E-39		5.0E-39	5.0E-39	4.0E-39	4.0E-39		4.0E-39	4.0E-39	4.0E-39	3.0E-39	3.0E-39
Expression Signal	1.41	0.83	1.31	1.31	1.21	3.61	3.61	2.78	6.23	2.33	5.14	1.51	9	25.5	2.12		2.23	1.3		5.13	1.54	10.53	96.0		1.49	3.08	2.03	11.27	11.27
ORF SEQ ID NO:	23743	23896		23903	24173	25682					19842			21832				20757		22682		20291	23238		26775				19831
Exen SEQ ID NO:	13965	14121	L	1_	14387	15582	1	Ι.		l	L.	<u></u>	L	11938	L	1	18337	10912		12884		10479	13441	L				Ш	10029
Probe SEQ ID NO:	4063	4223	4229	4229	4488	5673	5673	6378	7414	9264	8	1372	٤	2 2 8	8184		8838	99		2957	9556	538	3525		6707	9575	2896	4	4

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Table 4
Single Exon Probes Expressed in Heart

yd26g06.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:109402 5' similar to contains cocosa 10.s1 Soares_NhHMPu_S1 Homo sapiens cDNA clone IMAGE:1860988 3' similar to SW:GTR5_RAT P43427 GLUCOSE TRANSPORTER TYPE 5, SMALL INTESTINE; ox63a10.s1 Soares_NhHMPu_S1 Homo sapiens cDNA clone IMAGE:1660986 3' similar to SW:GTR5_RAT nw21g02.s1 NCI_CGAP_GCB0 Homo sapleris cDNA clone IMAGE:1241138 3' similar to contains THR.t3 zno6102.r1 Stratagene hNT neuron (#937233) Homo sapiens cDNA clone IMAGE:546651 5' finfc16 Regional genomic DNA specific cDNA library Homo sapiens cDNA clone CR12-1 yp51c06.s1 Soares retina N2b4HR Homo sapiens cDNA clone IMAGE:190954 3' 601301607F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3636289 5' Mus musculus mRNA for neuronal Interacting factor X 1 (NIX1) (NIX1 gene) Mus musculus mRNA for neuronal interacting factor X 1 (NIX1) (NIX1 gene) ng86f03.s1 NCI_CGAP_Pr6 Homo saplens cDNA clone IMAGE:941693 Homo saplens mRNA for ras-related GTP-biriding protein, complete cds Homo sapiens myosin phosphatase, target subunit 2 (MYPT2), mRNA Homo sapiens KVLQT1 gene P43427 GLUCOSE TRANSPORTER TYPE 5, SMALL INTESTINE Homo sapiens UDP-glucose pyrophosphorylase 2 (UGP2), mRNA Homo sapiens AE-binding protein 1 (AEBP1),mRNA Homo sapians homogantisata 1,2-dioxyganasa gane, completa cds PM0-BT0340-211299-003-402 BT0340 Homo sapians cDNA Homo sapiens DKFZp434P211 protein (DKFZP434P211), mRNA Homo sapiens DKFZp434P211 protein (DKFZP434P211), mRN **Top Hit Descriptor** EST384065 MAGE resequences, MAGB Homo saplens cDNA EST384065 MAGE resequences, MAGB Homo saplens cDNA Homo saplens chromosome 21 segment HS21C048 RC4-FN0037-280700-011-a10 FN0037 Homo saplens cDNA Alu repetitive element, contains LTR1 repetitive element; Homo saplens AE-blnding protein 1 (AEBP1) mRNA Homo sapiens tubby like protein 3 (TULP3), rnRNA promrna-7.D01.r bytumor Homo saplens cDNA Human mRNA for KIAA0209 gene, partial cds Homo sapiens KVLQT1 gene THR repetitive element; EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN **EST HUMAN EST HUMAN** HUMAN EST_HUMAN EST_HUMAN EST_HUMAN Top Hit Database Source HUMAN HUMAN EST EST 눋 눋 F 4755145 NT 11436738 NT 눋 z 7657020 NT 5803210 NT 4755145 7657020 11425464 Fop Hit Acesslon AW951995.1 .0E-39 AW951995.1 2.0E-39 AL163248.2 AA631949.1 AF000573.1 2.0E-39 AA720574.1 2.0E-39 BF370207.1 2.0E-39 AA080867.1 1.0E-39 AJ006345.1 1.0E-39 AJ278170.1 BE409203.1 AI525119.1 AW372318 AA508880. AJ006345.1 3.0E-39 AI084557.1 3.0E-39 AI084557.1 ģ D86964.1 1.0E-39 T80876.1 D78132.1 H37903.1 .0E-39 2.0E-39 (9.0E-40 9.0E-40 2.0E-39 9.0E-40 3.0E-39 2.0E-39 1.0E-39 1.0E-39 1.0E-39 3.0E-39 1.0E-39 Aost Similar (Top) Hit BLAST E Value 4.36 10.19 1.75 1.54 1.08 1.67 5.61 4.42 4.03 10.15 1.36 2.17 4.95 5.49 5.49 8.68 11.27 5.51 3.61 Expresslan Signal 20971 25142 25439 20294 ORF SÉQ ID NO: 28782 26384 21262 21275 24239 24240 24279 25460 25461 19832 28781 21707 21261 15395 15395 16028 11123 18900 10933 11418 15285 18587 14453 14453 15379 16220 10484 10029 18871 10820 12458 19394 11401 11401 11419 10805 14201 14492 18871 16223 SEQ ID 1216 6083 1215 4303 5459 6475 5475 Probe SEQ ID 9104 9104 9147 894 1015 1513 930 8860 9922 1497 1514 4561 4604 6357 543 4 879 1497 2587 4561 ö

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Top Hit Descriptor	Homo sapiens tissue inhibitor of metalloproteinase 3 (Sorsby fundus dystrophy, pseudoinflammatory) (TIMP3) mRNA	Homo saplens mRNA for KIAA1244 protein, partial cds	Homo sapiens ubiquitin specific protease 13 (isopeptidase T-3) (USP13) mRNA	Homo sapiens ubiquitin specific protease 13 (isopeptidase T-3) (USP13) mRNA	7H15A04 Chromosome 7 HeLa cDNA Library Homo sapiens cDNA clone 7H15A04	601288958F1 NIH_MGC_8 Homo sapiens cDNA clone IMAGE:3819166 5'	Human DNA polymerase gamma mRNA, nuclear gene encoding mitochondrial protein, complete cds	Human DNA polymerase gamma mRNA, nucléar gene encoding mitochondrial protein, complete cds	Homo sapiens chromosome 21 segment HS21,C046	EST70527 T-cell lymphoma Homo sapiens cDNA 6' end similar to similar to zinc finger protein family	EST70527 T-cell lymphoma Homo sapiens cDNA 5 end similar to similar to zinc finger protein family	hz40g01.x1 NCI_CGAP_GC6 Home sapiens cDNA clone IMAGE:3210480 3'	Homo sapiens fatty acid desaturase 1 (FADS1), mRNA	Homo sapiens fatty acid desaturase 1 (FADS1), mRNA	AV653028 GLC Homo sapiens cDNA clone GLCDGF04 3'	AV653028 GLC Hamo sapiens cDNA clane GLCDGF04 3'	Homo saplens chromosome 21 segment HS21,C085	tter bol x NCI_CGAP_Pr28 Homo sapiens cDNA clone IMAGE:2248873 3' similar to TR:073505 073505 POL PROTEIN.;	Homo sapiens X-linked anhidrollic ectodermal dysplasia protein gene (EDA), exon 2 and flanking repeat	regions	Homo sapiens KIAA0433 protein (KIAA0433), mRNA	nv34e10.r1 NCI_CGAP_Br4 Homo sapiens cDNA clone IMAGE:1222122	PM0-BN0167-070500-002-h12 BN0167 Homo saplens cDNA	PM0-BN0167-070500-002-h12 BN0167 Homc sapiens cDNA	RC1-CN0017-120200-012-e04 CN0017 Homo sapiens cDNA	wh12f07.x1 NCI_CGAP_Kid11 Homo saplens cDNA clone IMAGE:2380549 3'	Homo sepiens sema domain, seven thromboscondin repeats (type 1 and type 1-like), transmembrane domain (TM) and short cytoplasmic domain, (semaphorin) 5A (SEMA5A), mRNA
Top Hit Database Source	TN	NT	N	N	EST_HUMAN	EST_HUMAN	NT	IN	FZ	EST_HUMAN	EST HUMAN	EST HUMAN	N	LN	EST_HUMAN	EST_HUMAN	NT	EST_HUMAN		Ļ	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT
Top Hit Acession No.	4507512 NT	AB033070.1	4507848 NT	4507848 NT	AA078165.1	BE396541.1	J60325.1	J60325.1	AL163246.2	AA361275.1	4A361275.1	BE504766.1	11439783 NT	11439783 NT	AV653028.1	AV653028.1	AL163285.2	A1686005.1		AF003528.1	7682117 NT	AA742809.1	BE009416.1	BE009416.1	AW841585.1	AI825949.1	11417342 NT
Most Similar (Top) Hit BLAST E Value	9.0E-40	9.0E-40	9.0E-40	9.0E-40	8.0E-40	8.0E-40	7.0E-40	7.0E-40	7.0E-40	6.0E-40	6.0E-40	6.0E-40	6.0E-40	6.0E-40	6.0E-40	6.0E-40	5.0E-40	4.0E-40		4.0E-40	4.0E-40	4.0E-40	4.0E-40	4.0E-40	4.0E-40	3.0E-40	3.0E-40
Expression	5.04	3.58	0.82	1.12	0.95	2.41	1.56	1.58	2.83	3.88	3.88	2.07	3.08	3.08	6.82	6.82	1.89	1.38		2.67	7.85	3.76	4.87	4.87	4.07	0.98	6.25
ORF SEQ ID NO:				23923	22725		26578		28391	22449	22450		26124	26125		27805		21607			23973		27308			23721	26030
Exen SEQ ID NO:	11338	15069	14149	14149	12932	13758	16399	16399	18150	12560	12560	16531	15989	15989	17581	17581	12433	11731		11951	14189	16542	17114	17114	17989	13943	15906
Probe SEQ ID NO:	1433	3892	4250	4398	3004	3847	6541	6541	8270	2696	2696	5616	8141	6141	7731	7731	2561	1834		. 2861	4291	6662	7237	7237	8089	4040	6001

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SEQ Expression (Top) Hit Acession Database No. Signal BLAST E No. Source	26959 4.28 3.0E-40 5454167 NT Homo saplens HBV associated factor (XAP4) mRNA	1.49 3.0E-40 AF078779.1		19845 1 02 Contains MER29 bands septembly clone IMAGE:3146266 3' similar to contains MER29.b3	11.23 3.0E-40 6005813 NT	1 96 3 0E-40 AW118799 1 EST HUMAN	12.69 2.0E-40 AI223036.1 EST HUMAN	.1 EST_HUMAN	1.4 2.0E-40 AV731601.1 EST HUMAN AV731601 HTF Homo saplens cDNA clone HTFAZE05 5'		21664 2.16 2.0E-40 450618B NT products	Homo saplens proteasome (prosome, macropain) subunit, alpha type, 7 (PSMA7) mRNA, and translated products	21807 1.08 2.0E-40 AI968562.1 EST_HUMAN ZINC FINGER PROTEIN ;	2.81 2.0E-40 5453592	2.35		22808 3.59 2.0E-40 5453592 NT Homo sapiens adentyly cyclase-associated projein 2 (CAP2) mRNA	24478 1.49 2.0E-40 AL163280.2 NT Homo sapiens chromosome 21 segment HS21C080	2.0E-40 AL163280.2 NT		1.0E-40 BF036881.1	70,000,000	Т	0.02	1.0E-401BF-541030.1 EST HUMAN	1.81 1.0E-40 4507142 NT	6.28 1.0E-40 4508012 NT	24550 0.88 1.0E-40 7705778 NT Homo saplens CGI-65 protein (LOC51103) mRNA
							-	2.72	1.4							1.08				1.65								
SEQ ID ORF SEQ NO:	16761 2695	17063 2725	17188 2738	18284 2851				10701	11681		11787 2166	11787 2166	11917 2180	12011 2191		12525	13014 2280	14691 2447	14691 2447	10791	12451 2234	0000	12020					14772 2455
Probe E: SEQ ID SE(NO: N]	7186 1		8387	1		l	21 177	1783 1	l	1892 1	1892	2028 11	2123 13	1	2858 12	3087 13	4807 1		865 10		l	2020			- 1	4505 14	

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	_	_	_	_	_		_	_		_	_	_	_	_	_	_			L 48411	N P 11.5	_~		_		_		*****	****	
Top Hit Descriptor	nj42f04.s1 NCI_CGAP_AA1 Homo sapiens cDNA clone IMAGE:995167 3'	nj42f04.s1 NOI_CGAP_AA1 Homo sapiens cDNA clone IMAGE:995167 3'	AU149345 NT2RM4 Homo saplens cDNA clone NT2RM4002122 3'	qh31h10.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:1846339 3'	MR2-CT0222-211099-002-e10 CT0222 Homo sapiens cDNA	Homo saplens chromosome 21 segment HS21C003	wp04h04.x1 NCI_CGAP_Kid11 Hamo sapiens cDNA clone IMAGE:2463895 3'	wp04h04.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2463895 3'	Homo sapiens a disintegrin and metalloproteinese domain 22 (ADAM22), mRNA	Homo sapiens pescadilio (zebrafish) homolog 1, containing BRCT domain (PES1), mRNA	Homo sapiens DSCR5b mRNA, complete cds	Homo sapiens Down syndrome candidate regicn 1 (DSCR1), mRNA	601340485F1 NIH_MGC_53 Homo sapiens cDNA clone IMAGE:3682677 5	yc03e10.s1 Stratagene lung (#937210) Homo capiens cDNA clone IMAGE:796263'	Homo saplens target of myb1 (chicken) homolog (TOM1), mRNA	PM4-BT0341-251199-002-F11 BT0341 Homo sapiens cDNA	QV0-HT0367-150200-114-g09 HT0367 Homo; sapiens cDNA	AU119344 HEMBA1 Homo sapiens cDNA clone HEMBA1005583 5'	ow45e06.s1 Soares_parathyroid_tumor_NbHPA Homo sapiens cDNA clone IMAGE:1649794 3' similar to TR:000597 000597 CYTOCHROME C-LIKE POLYPEPTIDE. ;contains LTR5.b1 LTR5 repetitive element ;	ow45e06.s1 Soares_parathyroid_tumor_NbHPA Homo sapiens cDNA clone IMAGE:1649794 3' similar to TR:000597 000597 CYTOCHROME C-LIKE POLYPEPTIDE. ;contains LTR5.b1 LTR5 repetitive element ;	Homo sapiens gene for activin receptor type IIB, complete cds	tm96c04.x1 NCI_CGAP_Bm25 Home sapiens cDNA clone IMAGE:2165958 3' similar to contains OFR.b1 OFR repetitive element :	Homo sapiens 959 kb cantig between AML1 and CBR1 on chromosome 21q22; segment 1/3	Homo sapiens 959 kb contig between AML1 and CBR1 on chromosome 21q22; segment 1/3	H.sapiens DNase I hypersensitive site (HSS-3) enhancer element	AV758295 BM Homo saplens cDNA clone BMFBHC05 5'	601888098F1 NIH_MGC_17 Homo sapiens dDNA clone IMAGE:4122119 5'	AV710480 Cu Homo sapiens cDNA clone Cu/ACC07 5'	AV708431 ADC Homo sapiens cDNA clone ADCARE02 5'
Top Hit Database Source	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	- LN	EST_HUMAN	EST_HUMAN			N		EST_HUMAN	EST_HUMAN			EST_HUMAN	EST_HUMAN	EST_HUMAN		N	FST HIMAN	Т	LN.	NT	EST_HUMAN	EST_HUMAN		EST_HUMAN
Top Hit Acession No.	AA573201.1	AA573201.1	AU149345.1	AI239572.1	BF334112.1	AL163203.2	A1934364.1	AI934364.1	11419208 NT	11417972 NT	AB037163.1	7657042 NT	BE567816.1	T62628.1	4885636 NT	BE067042.1	BE156318.1	AU119344.1	A1027117.1	AI027117.1	AB008681.1	AIROADE 1	AJ229041.1	AJ229041.1		AV758295.1	BF304683.1	AV710480.1	AV708431.1
Most Similar (Top) Hit BLAST E Value	1.0E-40	1.0E-40/	1.0E-40 /	1.0E-40/	1.0E-40	8.0E-41	7.0E-41	7.0E-41	7.0E-41	7.0E-41	_	_	6.0E-41	5.0E-41	5.0E-41	5.0E-41	4.0E-41	4.0E-41	4.0E-41	4.0E-41	-	4 0F-41	-	4.0E-41	4.0E-41	4.0E-41	4.0E-41	_	4.0E-41
Expression Signal	2.03	2.03	5.72	53.3	3.93	1.73	2.36	2.36	3.27	4.82	1.72	2.19	0.94	1.31	0.98	2.55	1.91	1.28	9.42	9.42	2.12	α.	3.03	3.03	1.89	1.39	6.01	7.62	2.31
ORF SEQ ID NO:	26230	26231	28412	28480		26689			25663		20064	21851	24044	21532				20839	21149	21150		213BD		22579			27657		
Exen SEQ ID NO:	16081	16081	18168	18232	19622	16501	12678	12878	15567	19818	10244	11954	14260	11660	1	15850	10332	10998	11293	11293	1	41522	_	<u></u>	ļ	15825	ļ	18733	19510
Probe SEQ ID NO:	6215	6215	8289	8355	9521	6621	811	811	5655	9931	279	2064	4384	1761	4018	6945	385	1082	1388	1388	1404	1818	2859	2859	4052	5920	7590	8925	6696

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Table 4
Single Exon Probes Expressed in Heart

_				_																	**med			-50				····(· <i>)</i> ·	1171 #		***		1. "m."
	Top Hit Descriptor	Homo sapiens PAD-H19 mRNA for peptidylarginine deiminase type II, complete cds	Homo sapiens DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes, complete cds)	Homo sapiens mRNA for KIAA1327 protein, partial cds	H.sapiens mRNA for putative p64 CLOP protein	Homo saplens mRNA for KIAA1387 protein, partial cds	Human ribosomal protein L23a mRNA, complete cds	EST35818 Embryo, 8 week I Homo sapiens cDNA 5' end	Human mRNA for KIAA0207 gene, complete cds	G.gorilla DNA for ZNF80 gene homolog	Human ribosomal protein L23a mRNA, complete cds	Homo sapiens chromosome 21 segment HS21C067	Homo sapiens chromosome 21 segment HS21C067	Homo sapiens homolog of Nedd5 (hNedd5) mRNA, complete cds	Human B-cell specific transcription factor (BSAP) mRNA, complete cds	Human B-cell specific transcription factor (BSAP) mRNA, complete cds	EST31723 Embryo, 12 week I Homo sapiens cDNA 5' end	ZINC FINGER PROTEIN 135	EST84565 Colon adenocarcinoma IV Homo saplens cDNA 5' end	Mus musculus tubulin alpha 6 (Tuba6), mRNA	qf75c10.x1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1755858 3'	Homo sapiens hypothetical protein FLJ20454 (i ⁻ LJ20454), mRNA	RC0-HT0613-210300-032-g01 HT0613 Homo sapiens cDNA	Homo sapiens hypothetical C2H2 zinc finger protein FLJ22504 (FLJ22504), mRNA	Homo sapiens hypothetical C2H2 zinc finger protein FLJ22504 (FLJ22504), mRNA	Homo sapiens homeobox protein CDX4 (CDX4) gene, complete cds and flanking repeat regions	Homo saplens DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes,	complete cds)	nh07c02.s1 NCI_CGAP_Thy1 Homo sepiens cDNA clone IMAGE:943586 cimilar to TR:G434304 G434304	מסופר באר אביטובי לבינים ואל איינים א	Homo sapiens chromosome 21 segment HS21(C085	Homo sapiens phosphatidylinositol 4-kinase 230 (pI4K230) mRNA, complete cds	Homo sapiens phosphatidylinositol 4-kinase 230 (pi4K230) mRNA, complete cds
	Top Hit Database Source	N N	ΙN	N	LN	F	NT	EST_HUMAN	NT	NT	ΝΤ	FZ -	FN	ΙN	TN	LN FN	EST_HUMAN	SWISSPROT	EST_HUMAN	LN L	EST_HUMAN		EST_HUMAN	TN	LZ	IN.		. LN	MANUEL FOR	EST TOWAR	NT	FX	NT
	Top Hit Acession No.	4B030176.1	AB026898.1	3.0E-41 AB037748.1	X87689.1	3.0E-41 AB037808.1	J43701.1	.1	2.0E-41 D86962.1	2.0E-41 X89631.1	2.0E-41 U43701.1	4L163267.2	2.0E-41 AL163267.2	2.0E-41 AF038404.1	M96944.1	M96944.1	2.0E-41 AA328285.1	P52742	4A372637.1	6678468 NT	AI217868.1	11528291 NT	BE179191.1	11560151 NT	11560151 NT	AF003530.1		8.0E-42 AB026898.1	4 402000 4	8.UE-42 AA493690.1	L163285.2	F012872.1	AF012872.1
	Most Similar (Top) Hit BLAST E Value	3.0E-41 A	3.0E-41	3.0E-41	3.0E-41 X	3.0E-41	2.0E-41	2.0E-41	2.0E-41	2.0E-41				2.0E-41	2.0E-41 N	2.0E-41 N	2.0E-41	2.0E-41	2.0E-41	1.0E-41	1.0E-41	1.0E-41	9.0E-42 B	9.0E-42	9.0E-42	8.0E-42 A		8.0E-42	C7 30 0	8.UE-42			6.0E-42 A
	Expression Signal	<u>2</u> .	3.08	0.85	7.38	1.49	7.3	2.3	1.03	4.07	5.31	1.06	1.08	6.59	1.33	1.33	1.39	1.7	3.46	4.64	1.8	2.63	1.33	2.63	2.63	5.37		1.32	0	32.0	8.	3.25	3.25
	ORF SEQ ID NO:	20704	23914		25143	. 26873	21305	21688	21962	22010	21305	24189		26559	26772	26773	26786	27255	28882	24149	27494			27367	27368			21849				21586	
	Exan SEQ ID NO:	10857	14139	14919	15296	16765	11445	11810	12059	12108	11445	14414	14414	16381	16582	16582	16596	17065	18594	14359	17287	18930	16836	17168	L	10399	1	11952				L	11709
	Probe SEQ ID NO:	932	4240	5047	5376	6849	1782	1915	2172	2221	2798	4521	4521	6522	6702	6702	6716	7188	8777	4465	7420	9197	6958	7282	7292	455		2062	0000	8238	916	1812	1812

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Single Exon Probes Expressed in Heart

			_	-			-	-			T-1	6 1	_	$\overline{}$	-	~~	******		_	7			_	_			т	_	-	¬,
Top Hit Descriptor	xp29f08.x1 NCI_CGAP_HN10 Homo sapiens cDNA clone IMAGE:2741799 3' similar to contains L1.t1 L1 repetitive element ;	qu24h09.x1 NCI_CGAP_Br12 Homo sapiens cDNA clone IMAGE:1965761 similar to contains Alu repetitive element.	Homo saplens mRNA for KIAA1067 protein, pertial cds	Homo sapiens mRNA for KIAA1067 protein, partial cds	Homo sapiens Xq pseudoautosamal region; segment 1/2	hv31e11.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3175052 3'	Homo sapiens SET domain and mariner transposase fusion gene (SETMAR) mRNA	Homo sapiens SET domain and mariner transposase fusion gene (SETWAR) mRNA	Homo sapiens ubiquitin protein ligase E3A (human papilloma virus E6-associated protein, Angelman syndrome) (UBE3A), mRNA	Homo sapiens ubiquitin protein ligase E3A (human papilloma virus E6-associated protein, Angelmen syndrome) (UBE3A), mRNA	Homo saplens myctubularin related protein 3 (MTMR3), mRNA	Homo sapiens multifunctional calcium/calmodtilin-dependent protein kinase II detta2 isoform mRNA, complete	နောဘ	Homo sapiens mRNA for KIAA1294 protein, partial cds	Homo sapiens hypothetical protein FLJ20163 (FLJ20163), mRNA	Homo sapiens MHC class 1 region	Homo sapiens MHC class 1 region	Homo sapiens ribonuclease III (RN3) mRNA, complete cds	H.saplens PROS-27 mRNA	Homo sapiens regulatory factor X, 4 (influences HLA class II expression) (RFX4) mRNA	Homo sapiens zinc finger protein 177 (ZNF177) mRNA	RC1-ST0278-040400-018-h11 ST0278 Home sapiens cDNA	RC1-ST0278-040400-018-h11 ST0278 Home sapiens cDNA	601458531F1 NIH_MGC_66 Homo saptens cDNA clone IMAGE:3862086 5	ab14e10.s1 Stratagene lung (#937210) Homo sapiens cDNA clone IMAGE:840810 3' similar to contains	THR.12 THR repetitive element;	RC0-TN0079-110900-024-g07 TN0079 Homp sapiens cDNA	RC3-NN0070-270400-011-h10 NN0070 Homic sapiens cDNA	2819283.3prime NIH_MGC_7 Homo saplens cDNA clone IMAGE:2819293 3	EST387438 MAGE resequences, MAGC Horno sapiens cDNA
Top Hit Database Source	EST_HUMAN	EST HUMAN	NT	NT	IN	EST_HUMAN	TN	LN	FZ	L L	NT L		L	NT	ΝŢ	TN	NT	ΝT	Ŋ	IN	Ŋ	EST_HUMAN	EST_HUMAN	EST_HUMAN		EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN
Top Hit Acession No.	AW 238656.1	AI284770.1	AB028990.1	AB028990.1	AJ271735.1	BE217913.1	5730038 NT	5730038 NT	11433063 NT	11433083 NT	11417957 NT		AF071569.1	AB037715.1	8923162 NT	AF055068.1	AF055066.1	AF189011.1	X59417.1	4506496 NT	4508008 NT	AW818630.1	AW818630.1	BF035327.1		AA486105.1		AW698344.1		AW955368.1
Most Similar (Top) Hit BLAST E Value	6.0E-42	6 0F-42	_	_	5.0E-42	5.0E-42	5.0E-42	5.0E-42	5.0E-42	5 OF 42	6.0E-42		5.0E-42	5.0E-42	5.0E-42	4.0E-42	4.0E-42	4.0E-42	4.0E-42	4.0E-42	4.0E-42	4.0E-42				3.0E-42	2.0E-42	2.0E-42		2.0E-42
Expression Signal	3.51	20,	1.81	1.72	5.44	1.17	2.94	1.27	1.76	1 76	2.76		1.57	2.76	2.15	8.89	8.89	2.67	1.61	4.62	10.26	2.07	207	3.22		0.78	3.63	3.86	2.15	10.2
ORF SEQ ID NO:			25106			20197			26050		26203		26294		28494		20503	L			24162		L				21239			26500
SEQ ID NO:	12128			15275	10105	10376	10422	10423	•	1	ı		16138	16975	<u> </u>	L	10668	10987	L.	<u>_</u>	14374			_	L.	10084	L	12241	L	16437
Probe SEQ ID NO:	2244	4038	5355	5504	131	431	478	478	6018	8018	6072		6274	2098	8366	738	736	1050	4100	4156	4480	8041	8041	8714		86	1467	2361	2375	5519

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Top Hit Descriptor	EST367438 MAGE resequences, WAGC Homo sapiens cDNA	601081284F1 NIH_MGC_10 Homo sapiens cDNA clane IMAGE:3447620 5'	Human endogenous retrovirus pHE.1 (ERV9)	UI-H-BI1-afh-e-04-0-UI.s1 NCI_CGAP_Sub3 Homo sapiens cDNA clone IMAGE:2721871 3'	Homo sapiens partial C9 gene for complement component C9, exon 1	Homo sapiens partial C9 gene for complement component C9, exon 1	Homo sapiens NADH-ubiquinone oxdoreductase AGGG subunit precursor homolog mRNA, nuclear gene encoding mitochondrial protein, complete cds	Homo saplens NADH-ublquinone oxidoreductase AGGG subunit precursor homolog mRNA, nuclear gene	encoding mitochondrial protein, complete cds	Homo sapiens rec (LOC51201), mRNA	Homo sapiens PDNP1 gene, exon 17	Homo sapiens major histocompatibility complex, class II, DM alpha (HLA-DMA) mRNA	Homo sapiens origin recognition complex, subunit 5 (yeast homolog)-like (ORC5L) mRNA, and translated	products	Homo saplens KIAA0255 gene product (KIAA0255), mRNA	Homo sapiens chramosame 21 segment HS21,0087	Homo sapiens chromosome 21 segment HS21 C080	RC3-ST0197-161099-012-a03 ST0197 Homo saplens cDNA	Homo sapiens proteasome inhibitor (PI31), mRNA	Homo sapiens proteasame inhibitor (Pi31), mRNA	Homo saplens ryanodine receptor 3 (RYR3) mRNA	Homo sapiens mRNA for KIAA1288 protein, pertial cds	Homo sapiens a disintegrin and metalloproteinese domain 23 (ADAM23) mRNA	Homo saplens a disintegrin and metalloproteinese domain 23 (ADAM23) mRNA	Homo sapiens chromodomain protein, Y chromosome-like (CDYL) mRNA	z79a07.s1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:728532 3'	AV736824 CB Homo sapiens cDNA clone CBILAKH08 5	AV736824 CB Homo sapiens cDNA clone CBILAKH08 5'	Homo sapiens hypothetical protein FLJ20297 (FLJ20297), mRNA	Homo sapiens hypothetical protein FLJ20297 (FLJ20297), mRNA	Homo saplens hypothetical protein FLJ20297 (FLJ20297), mRNA	2822251.5prime NIH_MGC_7 Homo sapiens cDNA clone IMACE:2822251 6'
Top Hit Database Source	EST HUMAN	EST_HUMAN	LN	EST_HUMAN	TN	NT	LΝ		L	NT	NT	LN FA		NT	NT	ΙΝ	L'N	EST_HUMAN	N.	뉟	NT	. LN	NT	FN	TN	EST_HUMAN	HUMAN	EST_HUMAN	NT	ΙN		EST_HUMAN
Top Hit Acession No.		9.1	(57147.1	1.0E-42 AW 295809.1	1.0E-42 AJ251818.1	1.0E-42 AJ251818.1	F067168.1		F067166.1	11423219 NT	F110296.1	5174458 NT		4505524 NT	7662027 NT	1.0E-42 AL163267.2	1.0E-42 AL163280.2	1.0E-42 AW813817.1	5803122 NT	5803122 NT	4508758 NT	1.0E-42 AB033114.1	4501912 NT	4501912 NT	4757969 NT	VA435719.1	8.0E-43 AV736824.1	\V736824.1	8923276 NT	8923276 NT	8923276	W246442.1
Most Similar (Top) Hit BLAST E Value	2.0E-42	2.0E-42	1.0E-42 X57147.1	1.0E-42	1.0E-42	1.0E-42	1.0E-42		1.0E-42 A	1.0E-42	1.0E-42 A	1.0E-42		1.0E-42	1.0E-42	1.0E-42	1.0E-42	1.0E-42	1.0E-42	1.0E-42	1.0E-42	1.0E-42	1.0E-42	1.0E-42	9.0E-43	9.0E-43	8.0E-43	8.0E-43 A	8.0E-43			7.0E-43
Expression Signal	10.2			96.0	1.11	1.11	12.78		12.78	1.46	0.91	1.98		8.93	2.15	1.02	1.72	98.0	2.85	2.85	5.35	1.2	0.98	96.0	3.80	2.84	12.13	12.13	4.33	4.33	4.33	6.05
ORF SEQ ID NO:			20478	50789	20842	20843	20983		20984	21443	21772	22284		22662	23347	23553	23827	24170	24316	24317	24348	24442	24693	24694	27893	28525	20386	20387	20441	20442		23292
SEQ ID NO:			10648		11001	11001	12688	1	12688	11575	11879	12372				13760	14053	14383	14528	14528	14555	14654	14920	14920		L	10573	10573	10618	10618	10618	13503
Probe SEQ ID NO:	6219	7663	717	1026	1085	1085	1222		1222	1673	1986	2497		2935	3647	3849	4153	4489	4640	4840	4669	4769	. 5048	5048	7805	8397	636	මහ	685	685	685	3589

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Top Hit Descriptor	wp69b01.x1 NOI_CGAP_Brn25 Homo seplens cDNA clone IMAGE:2466985 3' similar to TR:015475 O15475 UNNAMED HERV-H PROTEIN ;contains LTR7.b1 LTR7 repetitive element;	ne72d06.s1 NCI_CGAP_Ew1 Homo sapiens cl2NA clone IMAGE:909803 similar to gb:L05095 60S RIBOSOMAL PROTEIN L30 (HUMAN);	AV708201 ADC Homo sapiens cDNA clone ADCACC10 5'	Homo saplens ATP-binding cassette, sub-family C (CFTR/MRP), member 3 (ABCC3), transcript variant MRP3B, mRNA	hd30b04.x1 Scares_NFL_T_GBC_S1 Hamo saplens cDNA clone IMAGE:2910991 3' similar to contains MER1.t3 MER1 MER1 repetitive element;	#35606.r1 Soares_NHHMPu_S1 Homo sapiens cDNA clone IMAGE:685410 5' similar to TR:G528641 G529641 DB1, COMPLETE CDS. ;contains elément PTR7 repetitive element ;	DKFZp761L1712_r1 761 (synonym: hamy2) Homo sapiens cDNA clone DKFZp761L1712 5'	Homo sapiens chromosome 21 segment HS21C013	EST86033 Testis I Homo sapiens cDNA 5' enc	П	Г	0052c10.x5 NCI_CGAP_Lu5 Homo sapiens cDNA clone IMAGE:1569810 3' similar to TR:P90591 P90591 PV14 GENE.;	DKFZp434D0119_r1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434D0119	MR2-SN0007-290400-004-c02 SN0007 Homo sapiens cDNA	55a4 Human retina cDNA randomly primed subilibrary Homo sapiens cDNA	Human mRNA for alpha-actinin	Homo sapiens X-linked anhidroitic ectodermal dysplasia protein gene (EDA), exon 2 and flanking repeat regions	Homo saplens protocadherin beta 6 (PCDHB6), mRNA	q178a02.x1 NCI_CGAP_Kid3 Homo sapiens d0NA clone IMAGE:1865354 3' similar to contains MER10.t3 MER10 repetitive element;	q76a02.x/1 NCI_CGAP_Kid3 Homo saplens cDNA clone IMAGE:1865354 3' similar to contains MER10.t3	MERTU repetitive element;	Value: 1 Sources retailing spiech in L.S. nome sapiens cultar done invade: 11 5027 3	yg0bb05.r1 Soares infant brain 1NIB Homo sapiens cDNA clone IMAGE:31363 5 similar to contains MEK10 repetitive element;	Homo sapiens calcium channel alpha1E suburit (CACNA1E) gens, exons 7-49, and partial cds, alternatively spliced
Top Hit Database Source	EST_HUMAN	EST HUMAN	EST_HUMAN	Ŋ	EST HUMAN	EST_HUMAN	EST_HUMAN	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	IN	LN	LN LN	EST HUMAN		ESI HUMAN	ESI HOMAN	EST_HUMAN	TN
Top Hit Acession No.	Al936748.1	6.0E-43 AA491890.1	6.0E-43 AV708201.1	8955973 NT	6.0E-43 AW 468897.1	6.0E-43 AA195164.1	6.0E-43 AL119158.1	5.0E-43 AL163213.2	5.0E-43 AA382780.1	5.0E-43 AV732578.1	5.0E-43 AA465288.1	5.0E-43 AI733244.1	AL049110.1	5.0E-43 AW863007.1	W29011.1	X15804.1	AF003528.1	11416793 NT	4.0E-43 AI244341.1		AI244341.1	17380.1	4.0E-43 R20950.1	AF223391.1
Most Similar (Top) Hit BLAST E Value	7.0E-43 A	6.0E-43				6.0E-43	6.0E-43	5.0E-43	6.0E-43	5.0E-43	5.0E-43	5.0E-43	5.0E-43 A	5.0E-43	5.0E-43 W.	5.0E-43	4.0E-43	4.0E-43	4.0E-43		4.0E-43 AI	4.05-43	4.0E-43	3.0E-43 A
Expression Signal	1.78	10.17	4.15	2.02	2.02	1.83	6.54	1.98	3.01	1.38	4.47	2.2	1.41	6.46	2.67	2.6	4.85	1.72	4.49		4.49	8.	1.89	2.84
ORF SEQ ID NO:				25829	28111	27745				22539	L					28146	20728		26824		26825			
Exan SEQ ID NO:	16869	11227	12421	15718	15975	17518	18322	10111	10437		17308	17795	17814			17802	12643		16636		L	- 0465	18915	11105
Probe SEQ ID NO:	7092	1320	2547	5811	6128	7668	8449	137	494	2816	7390	7945	7964	8145	8338	8753	957	6231	6757		9/2/	8024	9174	1195

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Single Exon Probes Expressed in Heart

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Top Hit Descriptor	H.sapiens gene encoding La autoantigen	AML1-EVI-1=AML1-EVI-1 fusion protein (rearranged translocation) [human, leukemic cell line SKH1, mRNA Mutant, 5938 nt]	nk66d06.s1 NCI CGAP Pr7 Homo sapiens cDNA clone IMAGE:1017419	Mus musculus otogelin (Otog), mRNA	Mus musculus ctogelin (Otog), mRNA	Human ribosomal RNA upstream binding transcription factor (UBTF) gene, partial cds	aa88111.s1 Stratagene fetal retina 937202 Horno sapiens cDNA clone IMAGE:838413 3' similar to contains THR.t2 THR repetitive element ;	Homo sapiens hypothetical protein (HSA011915), mRNA	Homo sapiens SET domain and mariner transposase fusion gene (SETMAR) mRNA	qd81c09.x1 Sogres_testis_NHT Homo septems cDNA clone IMAGE:1733968 3' similar to contains PTR7.t3 PTR7 PTR7 remaritius element	III.H.BIJdi-a-00.0111 st NOI CGAP Suk3 Homo seniens cDNA clone IMAGE 27247423'	Human ribosomal protein L23a mRNA, complete cds	FB1G5 Fetal brain. Stratagene Homo sapiens cDNA clone FB1G5 3'end similar to LINE-1	Homo sapiens Ras-like GTP-binding protein (FAB27A) gene, exons 1b and 2	Homo sapiens Ras-like GTP-binding protein (FAB27A) gene, exons 1b and 2	Homo sapiens chromosome 21 segment HS21(C084	602022313F1 NCI_CGAP_Brn67 Hamo sapiens cDNA clone IMAGE:4157666 5	Homo sapiens Sp4 transcription factor (SP4) niRNA	Homo sapiens Sp4 transcription factor (SP4) mRNA	vg40e01.r1 Soares infant brain 1NIB Homo sapiens cDNA clone IMAGE:34732 5' similar to SP:BD38 MOUSE P28656 BRAIN PROTEIN DN38 :	Homo sapiens 8q22.1 region and MTG8 (CBFA2T1) gene, partial cds	EST375749 MAGE resequences, MAGH Horrio saplens cDNA	wr87h01.x1 NCI_CGAP_Kid11 Homo saplens cDNA clone IMAGE:2494705 3'	Homo sapiens calcium channel, voltage-dependent, alpha 1E subunit (CACNA1E), mRNA	DKFZp761D1015_r1 761 (synonym: hamy2) Homo sapiens cDNA clone DKFZp761D1015 5'	wb99b04.x1 NCI_CGAP_Pr28 Homo sepiens oDNA clone IMAGE:2313776 3'	Homo sapiens cadherin EGF LAG seven-pass G-type receptor 1 (CELSR1), mRNA		qh23g01.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:1846552.3'	H.saplens DNA for Cone cGMP-PDE gene
Top Hit Database Source	N	Ļ	EST HUMAN	NT	LN PA	LN	EST_HUMAN	L	LN LN	EST HIMAN	ENT HIMAN	LO LO	EST HUMAN	Z	FN	L L	EST_HUMAN	LV	LN T	EST HUMAN	TN	EST_HUMAN	EST HUMAN	LN	EST_HUMAN	EST_HUMAN	TN	EST_HUMAN	EST_HUMAN	NT
Top Hit Acession No.	K97869.1	S69002.1	AA548154.1	7305360 NT	7305360 NT	J65487.1	44458824.1	7661721 NT	5730038 NT	A1100784 1	AW207390 4	143701.1	103007.1	AF154836.1	AF154836.1	AL 163284.2	BF348283.1	4507168 NT	4507168 NT	R19751.1	AF198490.1	AW963676.1	A1984961.1	11424378 NT	AL137964.1	AI675418.1	11418322 NT	AI222985.1	AI222985.1	X94354.1
Most Similar (Top) Hit BLAST E Value	3.0E-43	3.05-43	3.0E-43	3.0E-43	3.0E-43	3.0E-43	3.0E-43	3.0E-43	3.0E-43	2 OF 43	200	2 OF 43	2 0F-43	1.06-43	1.0E-43	1.0E-43	1.0E-43	1.0E-43	1.0E-43	1.0E-43	1.0E-43	1.0E-43	1.0E-43	1.0E-43	1.0E-43	1.0E-43	9.0E-44	8.0E-44	8.0E-44	8.0E-44
Expression Signal	1.48	1.05	0.95	1.71	1.71	3.78	6.68	1.18	2.02	4 27	1 28	7.83	338	2,92	2.92	3.36	4.95	12.07	12.07	.83	1.83	25.23	6.75	3.2	3.66	1.89	2.52	5.98	5.98	3.87
ORF SEQ ID NO:	21437					26072		27188			05590			21394	21395	21450	22444			24860		27199		28844		25311	25255	20648		27037
Exan SEQ ID NO:	11571	13440	14093	15743	15743	15940	16625	16997	18769	10148	18475	16715	18404	11534	11534	11580	12557	15892	15892	15117	16580	17006	18208	18560	18880	19058	19196	10798	10798	16845
Probe SEQ ID NO:	1669	3524	4193	5837	5837	6037	6746	7120	8962	14,	6343	6836	8532	1630	1630	1678	2692	5987	5987	6159	929	7129	8331	8872	9117	9405	9618	872	872	8889

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Top Hit Descriptor	Homo saplens mRNA for thymidine kinase, partial	Homo saplens myosin mRNA, partial cds	Homo sapiens polymerase (RNA) II (DNA directed) polypeptide F (POLR2F), mRNA	Homo sapiens putative nuclear protein (HRIHFB2122), mRNA	Homo saplens protein kinase C, alpha binding protein (PRKCABP), mRNA	Homo sapiens putative nuclear protein (HRIHFB2122), mRNA	ye89e01.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:124920 5'	Homo sapiens LIM domain-containing preferred translocation partner in lipoma (LPP) mRNA	Homo sapiens minisatellite ms32 repeat region	Homo sapiens minisatellite ms32 repeat region	Homo sapiens chromosome 21 segment HS21,0084	Homo saplens chromosome 21 unknown mRNA	Homo saplens chromosome 21 unknown mRNA	AU159839 Y79AA1 Homo saplens cDNA clone Y79AA1000496 3'	EST366120 MAGE resequences, MAGC Homo seplens cDNA	Homo saplens KIAA0851 gene (partial), XT3 gene and LZTFL1 gene	Homo sapiens KIAA0851 gene (partial), XT3 gene and LZTFL1 gene	bydodoz.x1 NCI_CGAP_Brn25 Homo sapiens cDNA clone IMAGE:2170083 3' similar to contains OFR.t1	AI 124571 NTORM4 Home septents oDNA clone NT2RM4000218 5	Homo sablens chromosome 21 segment HS21C103	Homo sapiens carboxyl terminal LIM domain protein (CLIM1) mRNA, complete cds	Homo sapiens karyopherin alpha 6 (importin alpha 7) (KPNA6), mRNA	601491529F1 NIH_MGC_69 Homo sapiens cDNA clone IMAGE:3893839 5'	zp18b05.r1 Stratagene fetal retina 937202 Horno sapiens cDNA clone IMAGE:609777 5'	Homo sapiens DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 1 (DDX1) mRNA	Homo sapiens DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 1 (DDX1) mRNA	Homo sapiens transmembrane trafficking protein (TMP21), mRNA	Homo sapiens transmembrane trafficking protein (TMP21), mRNA	Homo sapiens RAB36 (RAB36) mRNA, complete cds	hw14g08.x1 NCI_CGAP_Lu24 Homo sepiens\cDNA clone \MAGE:3182938 3' similar to SW:OXYB_HUMAN P22059 OXYSTEROL-BINDING PROTEIN .:	Homo sapiens tissue-type bone marrow zinc finger protein 4 mRNA, complete cds	Human mRNA for integrin alpha subunit, complete cds
Top Hit Database Source	N	NT	NT	NT	LZ		EST_HUMAN	NT	LN	NT	ΙN	LN LN	FZ	EST_HUMAN	EST_HUMAN	NT	LN	MANUT TOD	EST HIMAN	LN LN	TN	NT	T_HUMAN	EST_HUMAN	LN	NT	Z	N-	FN	FST HUMAN	NT	NT
Top Hit Acession No.	110498.2	29139.1	11527389 NT	11418088 NT	11418099 NT	11418088 NT	206035.1	5031886 NT	4F048729.1	3F048729.1	AL163284.2	4F231919.1	4F231919.1	AU159839.1	4W954050.1	4,1289880.1	4,1289880.1	4 000001	41306323.1	AI 183303 2	Igna7a 1	6912477 NT	3E880626.1	4A169851.1	4826685 NT	4826685 NT	5803200 NT	5803200 NT	4F133588.1	3F465325 4	AF070651.1	D25303.1
Most Similar (Top) Hit BLAST E Value	8.0E-44	8.0E-44	8.0E-44	8.0E-44	8.0E-44	8.0E-44	7.0E-44	7.0E-44	7.0E-44	7.05-44	7.0E-44	7.0E-44	7.0E-44	7.0E-44	6.0E-44	6.0E-44	6.0E-44	17	3.0E-44	A 0F 44	4 OF 44	3.0E-44	3.0E-44	3.0E-44	2.0E-44	2.0E-44	2.0E-44	2.0E-44	2.0E-44	2 OF 44	20E-44	2.0E-44
Expression Signal	3.39	5.06	2.59	1.89	1.75	1.84	0.85	12	22	2.2	2.28	1.17	1.17	2.05	2.51	2.52	2.04	0 10	3.60	200	12.2	1.07	45.1	5.08	2.75	2.75	4.63	4.63	2.79	1 43	2.03	1.1
ORF SEQ ID NO:	28643		25304	25191		25191		21978	22663	22664		23821	l		L				7/007	22006			22254	22777			L					22294
Exon SEQ ID NO:	18377	L	19031	19369	18532	19369	10580	12074	12864	12864	13698	14048	14048	16841	18791	10264	10288		16483			L	1			10951	L		L	t	Į.	Ιì
Probe SEQ ID NO:	8504	8935	9359	9400	9742	9885	643	2187	2837	2937	3788	4148	4148	6762	9868	38	328	1	2202	eec.	8584	1746	2485	3059	1033	1033	1189	1189	1290	1348	2105	2629

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Top Hit Descriptor	Homo sapiens adaptor-related protein complex 4, sigma 1 subunit (CLAPS4), mRNA	Homo sapiens DNA for amyloid precursor protein, complete cds	PM4-SN0016-120500-003-a04 SN0016 Homo saplens cDNA	Homo saplens chemokine (C-C motif) receptor 9 (CCR9), mRNA	Homo sapiens general transcription factor 2-I (GTF2I) mRNA, alternatively spliced product, complete cds	Homo sapiens glutamate receptor, metabotropic 3 (GRM3), mRNA	Homo sapiens glutamate receptor, metabotropic 3 (GRM3), mRNA	601286914F1 NIH_MGC_44 Homo seplens cDNA clone IMAGE:3613586 5'	TCBAP1E2795 Pediatric pre-B cell acute lymphoblastic leukemia Baylor-HGSC project=TCBA Homo sapiens cDNA clone TCBAP2795	Human mRNA for KIAA0376 gene, partial cds	Homo saplens cat eye syndrome chromosome region, candidate 1 (CECR1), mRNA	Homo sapiens Misshapen/NIK-related kinase (MINK), mRNA	Homo saplens Misshapen/NiK-related kinase (MINK), mRNA	RC1-CT0249-030300-028-h12 CT0249 Homo sapiens cDNA	RC1-BN0039-110300-012-b01 BN0039 Homo sapiens cDNA	Homo saplens chromosome 21 segment HS21C103	zw63d02.r/ Soares_total_fotus_Nb2HF8_9w Homo sapiens cDNA clone IMAGE:773763 5' similar to contains THR i3 THR repetitive element:	wested of the same total fature Nb2HF8 9w Home sablens cDNA clone IMAGE:773763 5' similar to	contains THR.t3 THR repetitive element;	zt88g11.r1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:728476 5'	Homo sapiens transcription factor IGHM enhancer 3, JM11 protein, JM4 protein, JM5 protein, T54 protein,	JM10 protein, A4 differentation-dependent protein, triple LIM domain protein 6, and synaptophysin genes,	Confidence cuts, and E-year cardinal areas	agotobusi soares innimiral of home sapiens cuna digne invacing a	Homo sapiens alpha satellite DNA, M1 monomer type	Homo sapiens alpha satellite DNA, M1 monomer type	AV714608 DCB Homo sapiens cDNA clone DCBBYE03 6'	Homo sapiens Sushi domain (SCR repeat) containing (BK65A8.2), mRNA	RC1-CT0198-150999-011-C08 CT0198 Homo sapiens cDNA	RC1-CT0198-150999-011-C08 CT0198 Homo sapiens cDNA	Homo sapiens hypothetical protein FLJ10379 (FLJ10379), mRNA
Top Hit Database Source	Z	NT	EST_HUMAN	NT	N.	LN	LΝ	EST_HUMAN	EST HUMAN	ΤN	Z-Z	ᅜ	N	EST_HUMAN	EST_HUMAN	ΙZ	EST HIMAN	AUMOLI - 10-	EST HUMAN	EST_HUMAN		ļ	101	EST_HUMAN	NT	INT	EST_HUMAN	NT	EST_HUMAN	EST_HUMAN	Ŋ
Top Hit Acession No.	5901933 NT	D87675.1	AW864379.1	11449901 NT	AF038968.1	11419226 NT	11419226 NT	BE389058.1	BE244902.1	AB002374.1	11526293 NT	7657334 NT	7657334 NT	AW853132.1	AW994803.1	AL163303.2	0.0434554.1	T. FOOTOTO	AA434554.1	AA398099.1			AF1807/8.1	AA455869.1	AJ130755.1	AJ130755.1	AV714608.1	10092664 NT	AW846967.1	AW846967.1	8922391 NT
Most Similar (Top) Hit BLAST E Value	2.0E-44	2.0E-44	2.0E-44	2.0E-44	2.0E-44	2.0E-44		2.0E-44	2.0E-44	2.0E-44	2.0E-44	1.0E-44	1.0E-44	1.0E-44	1.0E-44	1.0E-44	1 05 44	100	1.08-44	1.0E-44			1.0E-44	1.0E-44	1.0E-44	1.0E-44	1.0E-44	1.0E-44	1.0E-44		9.0E-45
Expression Signal	3.32	1.36	1.76	1.39	1.46	3.86	3.86	1.88	2.22	2.72	1.38	3.64	3.64	1.85	1.52	5.54	3 83	3	3.53	1.05		,	BE'L	5.08	.0.81	0.81	10.75	4.18	3.43	3.43	1.74
ORF SEQ ID NO:		23147	24152	25719	24870	26408	26407	27078		24910		19839	19840	20312			240gg	21300	21966				22489		24702	24703		28918			
Exan SEQ ID NO:	12438		14362	15617	16107	16245	16245	16886		I _	L		L_	L	I_	L	12054	1.	12064					13578	14931	14931	18255	18629	L.	18681	14370
Probe SEQ 10 NO:	2567	3425	4468	5709	6097	6383	6383	6007	9022	9710	9066	46	48	566	1179	1655	24.77	77	2177	2237			2732	3664	6061	5061	8378	8816	8869	8869	4476

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Top Hit Database Source	INT Homo saplens hypothetical protein FLJ10379 (FLJ10379), mRNA			au83h07x1 Schneider fetal brain 00004 Homo sapiens cDNA clone IMAGE:2782909 3' similar to	NOW T		T_HUMAN	(1994107.x1 NCI_CGAP_CLL1 Homo saplens cDNA clone IMAGE:2116453 3' similar to SW:PAX1_MOUSE_EST_HUMAN P09084 PAIRED BOX PROTEIN PAX-1.	Z172403.s1 Soares_testis_NHT Homo seplens cDNA clone IMAGE:727877 3' similar to contains element EST HUMAN TAR1 repetitive element;		BINT Homo sapiens golgin-like protein (GLP), mRNA	NT H.sepiens ART4 gene	HUMAN	EST_HUMAN		EST_HUMAN AV723976 HTB Homo saplens cDNA clone HTBAAG01 5	I/NT Homo sapiens golgi autoantigen, golgin subfanily a, 2 (GOLGA2) mRNA	NT Homo sapiens chromosome 21 segment HS2IC027	NT Homo sapiens chromosome 21 segment HS21C027	NT H. sapiens DNA for endogenous retroviral like element		NT Homo sapiens partial 5-HT4 receptor gene, exans 2 to 5	NT Hirmon ensinonhil Cherrott, I enden norstal (CLC) protein (heompsenholinsse) nane monarter and evon 1	T HUMAN	Г	aa87f12.r1 Stratagene fetal retina 937202 Homio sapiens cDNA clone IMAGE:838319 5' sImilar to EST HUMAN TR:G1144569 G1144569 R-SLY1.	Ī		
Top Hit Acession No.	8922391 NT	5174718 NT	5174718 NT	034167670.4		AL 1632		AI523768.1	AA397781.1		8923698 NT	X95826.1		BF676077.1	T71480.1	AV723976.1	4758451	-			AL163218.2	AJ243213.1) 01685 4						
Most Similar (Top) Hit BLAST E Value	9.0E-45	8.0E-45	8.0E-45	90 30	8.0E-43	5.0E-45	5.0E-45	5.0E-45	5.0E-45	5.0E-45	6.0E-46	4.0E-45	4.0E-45	4.0E-45	3.0E-45	3.0E-45	3.0E-45	3.0E-45	3.0E-45	3.0E-45	2.0E-45	2.0E-45	2 OF 45	2.0E-45	2.0E-48	2.0E-45	2.0E-45	2.0E-45	2 0F 45
Expression Signal	1.74	3.9	7.49	30.3	0.50	1.1	5.01	2.01	8.83	1.67	2.67	8.96	1.98	1.62	1.17	1.51	3.44	8.37	8.37	1.33	2.17	0.93	4 80	1.75	27.64	3.96	2.13	2.13	276
ORF SEQ ID NO:	24160	22245					21739	22904	25155	27292	29043	20883	22027				27173	27996	27997			22716	25057	26518	28315	28652			
Exon SEQ ID NO:	14370	12353	14889	13808			L	13098	15303	17103	18748	11041	12130	19192	13210		16981	17757	17757	19660	12331	12924	45834		Ĺ	18388	L		L
Probe SEQ ID NO:	4478	2477	5015	2002	0707	874	1957	3173	5384	7226	8940	1127	2248	9613	3997	6920	7104	7907	7807	9814	2454	2998	5020	6489	8179	8516	8794	8794	ORGS

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Probe SEQ ID NO:	SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
118	10349		2.22	1.0E-45	BE389855.1	EST_HUMAN	601284360F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3608183 5'
403	10349		2.7	1.0E-45	BE389855.1	EST_HUMAN	601284360F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3606183 5'
464	10407	20227	1.5	1.0E-45	4506412 NT	IN	Homo sapiens RAP1A, member of RAS oncogene family (RAP1A), mRNA
1157	11070	20915	1.1	1.0E-45	IN 0627280	NT	Homo sapiens Langerhans cell specific c-type lectin (LANGERIN), mRNA
3065	12992	22784	92'9	1.0E-45	U32169.1	IN	Human pro-a2 chain of collagen type XI (COL1/1A2) gene, complete cds
3447	13384	23171	1.04	1.0E-45	IN 8556598	IN	Homo saplens chromosome 21 open reading frame 1 (C21orf4), mRNA
3526	13442	23239	0.81	1.0E-45	AB046811.1	N	Homo sapiens mRNA for KIAA1591 protein, partial cds
4378	14274	24055	4.08	1.0E-45	BE396633.1	EST_HUMAN	601289116F1 NIH_MGC_B Homo sapiens cDINA clone IMAGE:3619803 5'
4884	14765	24541	1.05	1.0E-45	11545796 NT	NT	Homo sapiens niban protein (NIBAN), mRNA
7274	17151	27347	5.22	1.0E-45	BE887843.1	EST_HUMAN	601511226F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3912535 5"
7485	17355	27559	1.25	1.0E-45	AB002297.1	NT	Human mRNA for KIAA0299 gene, partial cds
9231	18950	25358	4.3	1.0E-45	TN 66081411	IN	Homo saplens protein kinase C, alpha binding protein (PRKCABP), mRNA
9415	19067		5.38	1.0E-45	11526291 NT	NT	Homo sapiens hypothetical protein FLJ20454 (FLJ20454), mRNA
9421	19070		2.58	1.0E-45	11418177 NT	L	Homo sapiens Ran GTPase activating protein 1 (RANGAP1), mRNA
9820			3.17	1.0E-45	11418157 NT	NT	Homo saplens calcium channel, voltage-dependent, alpha 11 subunit (CACNA11), mRNA
6790	16669	26861	2.28	9.0E-46	TN 6620166	. IN	Mus musculus karatin complex 2, gene 6g (Krt2-6g), mRNA
7018	16895		6.71	9.0E-48	AL163209.2	IN	Homo saplens chromosome 21 segment HS21C009
7986	17836	28077	7.89	9.0E-46	AW246964.1	EST_HUMAN	2822449.5prime NIH_MGC_7 Homo sapiens cDNA clone IMAGE:2822449 5'
0300	02000		9			MALA TOD	132/08.x1 NCI_CGAP_Gas4 Homo sapiens ciDNA clone IMAGE:2132189 3' similar to gb:J00314_ma2
7807	L	7	0.0	0.0E-40	A1455201.1	NAMOR 163	IODOLIN DELATIONALIA),
2392	12270	22166	8.79	8.0E-46	A1433261.1	EST_HUMAN	t32/08.x1 NCI_CGAP_Gas4 Homo saplens cONA clone IMAGE:2132199 3' similar to gb:J00314_ma2 TUBULIN BETA-1 CHAIN (HUMAN);
6692	16572		3.97	8.0E-48	BE167244.1	EST_HUMAN	RC5-HT0506-280200-012-C12 HT0508 Homo sapiens cDNA
4478	14372		6.54	7.0E-48	BE386165.1	EST_HUMAN	601277292F1 NIH_MGC_20 Homo sapiens cDNA clone IMAGE:3618119 5'
4701	14587		1.01	7.0E-48	BE084388.1	EST_HUMAN	RC4-BT0310-110300-015-f10 BT0310 Homo saplens cDNA
5683	15592		4.01	7.0E-46	TN 8922268	LN L	Homo sapiens hypothetical protein FLJ10847 (FLJ10847), mRNA
5912	15818	25943	1.35	7.0E-48	BF105845.1	EST_HUMAN	601822835F1 NIH_MGC_77 Homo sapiens cDNA clone IMAGE:4042736 5'
9543	19144		1.35	7.0E-46	AL163246.2	NT	Homo sapiens chromosome 21 segment HS21C046
							wm31f08.x1 NCI_CGAP_Ut4 Homo seplens cDNA clone IMAGE:2437575 3' similar to contains MER19.t2
2726	12588	22483	5.53	6.0E-46	AI884381.1	EST_HUMAN	MER19 repetitive element;
	L						wm31f08x1 NCI_CGAP_Ut4 Homo saplens cDNA done IMAGE.2437575 3' similar to contains MER19.t2
2726	12588	22484	5,53	6.0E-46	AI884381.1	EST_HUMAN	MER19 repetitive element;
5727	15634	25737	8.85	6.0E-46	A1635448.1	EST HUMAN	ts58h10.x1 NCI_CGAP_Kid8 Homo sepiens cDNA clone IMAGE:2232835 3' similar to TR:060363 060363 SA GENE. :
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Top Hit Descriptor	601478409F1 NIH_MGC_68 Homo sapiens cDNA clone IMAGE:3880895 5'	Homo sapiens chromosome 21 segment HS21C010	7d81g01.x1 Lupski_dorsal_root_ganglion Homo sapiens cDNA clone IMAGE:3279408 3'	7d81g01.x1 Lupski_dorsal_root_ganglion Home sapiens cDNA clone IMAGE:3279408.3	nea38f07.x1 NCI_CGAP_Kid11 Homo sapiens dDNA clone IMAGE:3258757 3' similar to TR:075202 075202 HOMOLOG OF RAT KIDNEY-SPECIFIC;	602021164F1 NCI_CGAP_Brn67 Homo sapiens cDNA clone IMAGE:4156670 5'	no54e09.s1 NCI_CGAP_SS1 Homo sapiens cDNA clone IMAGE:1104520 3' similar to gb:X53741_rna1 FIBULIN-1, ISOFORM A PRECURSOR (HUMAN);	hi86c03.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3008B36 3' similar to gb:X14008_rns1 LYSOZYME C PRECURSOR (HUMAN);contgins element MER37 repetitive element ;	hl88c03.x1 NC_CGAP_Lu24 Homo saplens cDNA clone IMAGE:3008838 3' similar to gb:X14008_rna1 LYSOZYME C PRECURSOR (HUMAN);contains element MER37 repetitive element;	Human endogenous retrovirus RTVL-H2	Homo sapiens mRNA for KIAA0622 protein, partial cds	Homo sapiens mRNA for KIAA0622 protein, partial cds	Human ig germilne gamma-3 heavy-chain gene V region, partial cds	Human Ig germline gamme-3 heavy-chain gene V region, partial cds	Homo sapiens DNA for Human P2XM, complete cds	Homo saplens mitogen-activated protein kinase kinase kinase 3 (MAP4K3), mRNA	H.sapiens Ig lambda light chain varlable region gene (7o.11.2) germline; Ig-Light-Lambda; VLambda	H.sapiens Ig lambda light chain variable region gene (7c.11.2) germline, Ig-Light-Lambda; VLambda	wj49c04.x1 NCI_CGAP_Lu19 Homo sapiens cDNA clone IMAGE:2406150 3' similar to contains THR.b2 THR repetitive element ;	Human mRNA for KIAA0061 gene, partial cds	ne08a09.s1 NCI_CGAP_Co3 Homo sapiens cDNA clone IMAGE:880408 3' similar to contains THR.b2 THR repetitive element;	Z27a11.s1 Soares_fetal_liver_spleen_1NFLS_IS1 Homo sapiens cDNA clone IMAGE:431996 3'	Homo sapiens Bruton's tyrosine kinese (BTK), alpha-D-galactosidase A (GLA), L44-like ribosomal protein (L44L) and FTP3 (FTP3) genes, complete cds
Top Hit Database Source	EST_HUMAN	LN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	١	Ŋ	TN	N	NT	TN	TN	Ę	EST_HUMAN	Z-L	EST_HUMAN	EST_HUMAN	NT
Top Hit Acession No.	BE784971.1	AL163210.2	BE677194.1	BE677194.1	BF590442.1	BF347229.1	AA601143.1	AW770544.1	AW 770544.1	M18048.1	AB014522.1	AB014522.1	M36852.1	M36852.1	AB002059.1	4506376 NT	273660.1	273660.1	AI831462.1	D31785.1	AA488646.1	AA678246.1	U78027.1
Most Similar (Top) Hit BLAST E Value	6.0E-46	5.0E-48		5.0E-46	5.0E-46	5.0E-46	4.0E-46	4.0E-48	4.0E-48	4.0E-48	4.0E-48	4.0E-48	4.0E-46	4.0E-46	4.0E-46	3.0E-46	3.0E-46	3.0E-46	3.0E-48	3.0E-48		2.0E-48	
Expression Signal	3.03	9.41	1.12	1.12	1.79	3.52	1.51	3.57	3.57	3.55	1.07	1.07	1.84	1.84	1.91	8.0	1.13	1.13	8.3	2.63	5.91	1.32	2.43
ORF SEQ ID NO:			23205	23206	26074			21446	21447	22464	23999	24000	25082	25083	25237	92662	24343	24344	27151		, 20594		21385
Exan SEQ ID NO:	17878	10169	13400	13400	15942	16017	10562	11578	11578	12573		14217	15258	L	19221	14192	14552	14552	16958	18668	10747	11447	
Probe SEQ ID NO:	8694	197	3484	3484	6039	6144	625	1676	1676	2710	4320	4320	5338	5338	0996	4584	4665	4666	7081	8856	819	1542	1623

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Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
4899	14779	24555	1.15		AA399286.1	EST_HUMAN	259902.r1 Soares, testis, NHT Homo sapiens cDNA clone IMAGE:726650 5' similar to SW:RSP1_MOUSE Q01730 RSP-1 PROTEIN ;
6418	16280	26442	6.78	2.0E-46	9910569 NT	IN	Mus musculus sperm tail associated protein (Stap), mRNA
6703	16583		1.17		BE86915	EST_HUMAN	601445137F1 NIH_MGC_65 Homo sapiens cDNA clone IMAGE:3849297 5'
8571	18439		1.87	2.0E-46	7657233 NT	LΝ	Homo sapiens small acidic protein (IMAGE145052), mRNA
9157	19629		1.75		BF028854.1	EST_HUMAN	801765225F1 NIH_MGC_53 Homo sapiens cDNA clone IMAGE:3997326 5'
8408	19525		1.44		H48391.1	EST_HUMAN	Jr32d01.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:206977 5'
9728	19515	25136	3.81	2.0E-46	AW277214.1	EST_HUMAN	xq78h03.x1 NOL_CGAP_Lu34 Homo saplens cDNA clone IMAGE:2758789 3'
1213	11121	20970	5.19	1.0E-46	4502694 NT	۲	Homo sapiens cell division cycle 10 (homologous to CDC10 of S. cerevisiae) (CDC10) mRNA
2236		22023	4.6	1.0E-48	AW978516.1	EST_HUMAN	EST390625 MAGE resequences, MAGP Homo sapiens cDNA
2351	12231	22128	2.59	1.0E-48	H97330.1	EST_HUMAN	EST48b095 WATM1 Homo saplens cDNA clone 48b095
3211	13135	22936	281	1 05-46	AA631912.1	HST HIMAN	np78b02.s1 NCI_CGAP_Pr2 Homo sapiens cDNA done IMAGE:1132395 similar to gb:X76717 H.sapiens MT-11 mRNA (HUMAN):
4772	1_			L	AB023197.1	L L	Homo sapiens mRNA for KIAA0980 protein, partial cds
5495		25477	4.18		BF194707.1	EST HUMAN	7692b01.x1 NCI_CGAP_Ov18 Homo sepiens cDNA clone IMAGE:3843705 3'
5636	1	25639	5.66	1.0E-46	8923762 NT	NT	Homo sapiens centaurin-alpha 2 protein (HSA272185), mRNA
5636		25640	5.66	1.0E-46	8923762 NT	N	Homo saplens centaurin-alpha 2 protein (HSA272195), mRNA
8236		25477	4.26		BF194707.1	EST_HUMAN	7092b01.x1 NCI_CGAP_Ov18 Homo sapiens cDNA clone IMAGE:3643705 3'
9188	18923		1.43	1.0E-46	BF531102.1	EST_HUMAN	602072264F1 NCI_CGAP_Brn67 Horno saplens cDNA clone IMAGE:4215398 5'
9188	1	25349		1.0E-48	BF531102.1	EST_HUMAN	602072264F1 NCI_CGAP_Brn67 Homo saplens cDNA clone IMAGE:4215398 5'
9925	19397		1.53	1.0E-46	AV715377.1	EST_HUMAN	AV715377 DCB Homo sapiens cDNA clone DCBAIE03 5'
749	10679		4.51	9.0E-47	AJ271735.1	IN	Homo saplens Xq pseudoautosomal region; segment 1/2
4848	14729	24512	197	9.0F-47	AW770928.1	FST HUMAN	hig3e04.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clane IMAGE:3009534 3' similar to TR:075703 075703 HYPOTHETICAL 12.4 KD PROTEIN
9682					11417966 NT	NT	Homo saplens SEC14 (S. cerevisiae)-like 2 (SEC14L2), mRNA
1768	11665		14.02	8.0E-47	Y18536.1	Z-L	Homo sapiens HLA-C gene, exon 5, individual 19323
1766	11665		14.02		Y18536.1	NT	Homo sapiens HLA-C gene, exon 5, individual 19323
2684	12549	22439	1.74	8.0E-47	5453956 NT ::	: LN	Homo sapiens protein phosphatase 2, regulatory subunit B (B56), epsilon isoform (PPP2R5E) mRNA
2989	12917		1.72	8.0E-47	AJ229043.1	LN LN	Homo sapiens 959 kb contig between AML1 and CBR1 on chromosome 21q22, segment 3/3
2501	12376		3.05	6.0E-47	AL163246.2	LΝ	Homo sapiens chromosome 21 segment HS21 C046
7344			6.27		A1695189.1	EST_HUMAN	tz98h02.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2296659 3'
5963		25990	5.27		11423972 NT	NT	Homo sapiens CDC37 (cell division cycle 37, S. cerevisiae, homolog) (CDC37), mRNA
8174	18062		3.91	5.0E-47	M78590.1	EST_HUMAN	EST00738 Fetal brain, Strategene (cat#936203) Homo sapiens cDNA clone HFBCF07

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Table 4
Single Exon Probes Expressed in Heart

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Top Hit Descriptor	Homo sapiens E1A binding protein p300 (EP300) mRNA	601280486F1 NIH_MGC_39 Homo sapiens cDNA clone IMAGE:3622437 5'	601280486F1 NIH_MGC_39 Homo sapiens cDNA clone IMAGE:3622437 5	xx66b07.x1 NCI_CGAP_Lym12 Homo sapienė, cDNA clone IMAGE:2848597 3' similar to SW:INT6_MOUSE. Q64252 VIRAL INTEGRATION SITE PROTEIN INT-6. [1];	601497639F1 NIH_MGC_70 Homo sapiens clJNA clone IMAGE:3899721 5'	601497639F1 NIH_MGC_70 Homo sapiens cDNA clane IMAGE:3898721 5'	yy54b04.s1 Soares_multiple_scierosis_2NbHMSP Homo sapiens cDNA clone IMAGE:277327 3	Homo sapiens chromosome 21 segment HS21C084	Homo sapiens KIAA0439 mRNA, partial cds	Homo sapiens nuclear dual-specificity phosphatase (SBF1) mRNA, partial cds	Human T-cell receptor active alphe-chain mRNA from JM cell line, complete cds	UI-HF-BM0-adx-d-07-0-UI.r1 NIH_MGC_38 Homo sapiens cDNA clane IMAGE:3063205 5'	UI-HF-BM0-adx-d-07-0-UI.r1 NIH_MGC_38 Homo sapiens cDNA clone IMAGE:3063206 5	qh04e07.x1 Soares_NFL_T_GBC_S1 Home saplens cDNA clone IMAGE:1843716 3'	Homo sapiens myosin phosphatase, target subunit 2 (MYPT2), mRNA	Homo saplens chromosome 21 segment HS2; C009	Homo sapiens chromosome 21 segment HS270009	wq98b02.x1 NCI_CGAP_GC8 Homo sapiens cDNA clone IMAGE:2479851 3'	Homo sapiens KIAA0426 gene product (KIAA0426), mRNA	ng43h12.s1 NCI_CGAP_Co3 Homo saplens cDNA clone IMAGE:937607 3'	Homo sapiens promyelocytic leukemia zlnc fingar protein (PLZF) gene, complete cds	Homo sapiens ring finger protein (C3HC4 type) 8 (RNF8), mRNA	nf23g07.s1 NCI_CGAP_Pr1 Homo saplens dDNA clone IMAGE:914652	nf23g07.s1 NCI_CGAP_Pr1 Homo saplens cDNA clone IMAGE:914652	Homo sapiens Rev/Rex activation domain binding protein-related (RAB-R) mRNA	EST377239 MAGE resequences, MAGI Homo sapiens cDNA	601463932F1 NIH_MGC_67 Homo sapiens cDNA clone IMAGE:3867487 5'	601463932F1 NIH_MGC_67 Homo sapiens cDNA clone IMAGE:3867487 5	Homo saplens 5-hydroxytryptamine 1D receptor pseudogene with an Alu repeat insertion	Homo sapiens DNA for amyloid precursor protein, complete cds	Homo saplens DNA for amyloid precursor projein, complete cds	Homo saplens SPH-binding factor mRNA, partial cds	Homo sapiens myosin phosphatase, target subunit 2 (MYPT2), mRNA
Top Hit Database Source		EST_HUMAN	EST_HUMAN	EST_HUMAN			T_HUMAN	NT	NT		LN	EST_HUMAN	EST_HUMAN	EST_HUMAN	FX	IN	١	EST_HUMAN		EST_HUMAN	NT	NT	EST_HUMAN	EST_HUMAN	TN	EST_HUMAN	EST_HUMAN	EST_HUMAN	IN	된	NT	NT	INT
Top Hit Acession No.	4557556 NT	BE616483.1	BE616483.1	AW 515509.1	BE807634.1	BE907634.1	N57483.1	AL163284.2	AB007899.1	U93181.1	M12959.1	AW 408800.1	AW 408800.1	AI222413.1	4505318 NT	AL163209.2	AL163209.2	Al969279.1	7662109 NT	AA524514.1	AF060568.1	4504866 NT	AA569592.1	AA56959	5174648 NT	AW8651	_	BE778475.1	L09731.1	D87675.1	D87675.1	AF07177	4505318 NT
Most Similar (Top) Hit BLAST E Value	4.0E-47	4.0E-47	4.0E-47	4.0E-47	3.0E-47	3.0E-47	3.0E-47	3.0E-47	3.05-47	3.0E-47	3.0E-47	3.0E-47	3.0E-47	3.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47
Expression Signal	3.41	2.08	2.08	4.84	1.75	1.75	6.17	8.9	1.5	4.99	76.0	4.26	4.26	1.89	4.27	2.18	2.18	1.18	1.22	4.44	2.3	1.66	1.76	1.75	2.06	1.1	1.6	1.6	1.33	2.1	2.1	1.77	5.77
ORF SEQ ID NO:	21140				20287	20288		20703	21770		23941				19936				21338	١.			L			L			L	28716		L	19936
Exen SEQ ID NO:	11284	16816	16816	18713	10474	10474	10730	10855	11877	13796	14164	15571	15571	15858	10118	10876	10876		L	L.		L	14186	14186	L.	L	L	١.				ı	1
Probe SEQ ID NO:	1379	6938	8669	8905	532	532	8	830	1984	3885	4265	5659	6999	5953	142	952	952	1548	1676	1654	2185	4261	4287	4287	4408	4707	5635	5635	6532	6643	6643	7060	9070

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Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acessian No.	Top Hit Database Source	Top Hit Descriptor
9219	19658	24988	1.98	2.0E-47	R42423.1	EST_HUMAN	yf92e08.s1 Soares infant brain 1NIB Homo sepiens cDNA clone IMAGE:29966 3' similar to contains OFR repetitive element;
9257	19659		1.32	2.0E-47	2.0E-47 AL163209.2	NT	Homo saplens chromosome 21 segment HS21009
1383	11288	21142	4.5	1.0E-47	Al333429.1	EST_HUMAN	qp99h03.x1 Soares_fetal_lung_NbHL19W Homo sapiens cDNA clone IMAGE:1931189 3'
3749	13662	23443	62'0	1.0E-47	BE280477.1	EST_HUMAN	601155321F1 NIH_MGC_21 Homo sapiens cDNA done IMAGE:3138893 5'
3749	13662	23444	67.0	1	1.0E-47 BE280477.1	EST_HUMAN	601155321F1 NIH_MGC_21 Homo saplens cDNA done IMAGE:3138893 5'
5008	14882		2.59	1.0E-47	1.0E-47 AW813906.1	EST_HUMAN	RC3-ST0197-130400-017-h02 ST0187 Homo sapiens cDNA
6194	15954	26086	7.68	1.0Ė-47 AI	A1880886.1	EST HUMAN	at19e08.x1 Barstead aorta HPLRB9 Homo sapiens cDNA clone IMAGE:2355686 3' similar to gb:M22995 RAS-RELATED PROTEIN RAP-1A (HUMAN)
7928	17778	28017		1.0E-47	10115.1	NT	Papio hamadryas alcohol dehydrogenase class (ADH) gene, 5' region
1605	11400	213EB	234	9 OF 48	AF223391 1	I-Z	Homo sapiens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively spliced
3509	1	23228			9.0E-48 BF359947.1	EST HUMAN	CM2-MT0100-310700-290-f05 MT0100 Homo sapiens cDNA
8460	1				Γ	EST HUMAN	601310479F1 NIH_MGC_44 Homo sapiens cDNA clone IMACE:3632083 5
1230	11138		1.32		18	1.	Homo sapiens aminoacylase 1 (ACY1), mRNA
1231	11138		1.51	8.0E-48	4501900 NT	N _T	Homo sapiens aminoacylase 1 (ACY1), mRNA
3098	13023	22818	3.62	8.0E-48 A	AW 768477.1	EST_HUMAN	hk61b03.x1 NCI_CGAP_Lym12 Homo sapiens cDNA clone IMAGE:3001133 3' similar to gb:X64707 BREAST BASIC CONSERVED PROTEIN 1 (HUMAN);
3098	13023	22819	3.62	8.0E-48	8.0E-48 AW 768477.1	EST HUMAN	hk61b03.x1 NCI_CGAP_Lym12 Homo sapiens cDNA clone IMAGE:3001133 3' similar to gb:X64707 BREAST BASIC CONSERVED PROTEIN 1 (HUMAN);
482				7.0E-48		NT	Homo sapiens mRNA for KIAA1209 protein, pertial cds
483	10426		13.37	7.0E-48	7.0E-48 AB033035.1	TN	Homo sapiens mRNA for KIAA1209 protein, partial cds
1482	11387	21260	1.12	7.0E-48	TN 6912719	NT	Homo sapiens tousled-like kinase 1 (TLK1), mRNA
1620	11524	21382	3.49	7.0E-48	IN 8600678	NT	Homo sapiens SET domain and mariner transposase fusion gene (SETMAR) mRNA
5947	15852		22.88	7.0E-48	11416831 NT	ĮN	Homo saplens histidy/-fRNA synthetase (HARS), mRNA
7275	17152		1.62	6.0E-48	6.0E-48 AF026816.1	. LN	Homo sapiens putative oncogene protein mRNA, partial cds
7500		27577	1.9		11427428 NT	NT	Homo sapiens hypothetical protein FLJ11006 (FLJ11006), mRNA
							zq45b06.s1 Stratagene hNT neuron (#937233) Homo saplens cDNA clone IMAGE:632627 3' similar to
7587	17438	27654		6.0E-48	AA18908	EST_HUMAN	contains Alu repetitive element;
3269				6.0E-48	4826891 NT	NT	Homo sapiens phosphodiesterase 1A, calmodulIn-dependent (PDE1A) mRNA
8325	18202		3.55	4.0E-48	4.0E-48 AI620420.1	EST_HUMAN	tu47a02.x1 NCI_CGAP_Pr28 Homo sapiens cDNA clone IMAGE:2254154 3'
1363			0.92	3.0E-48 A	V69096	EST_HUMAN	AV690964 GKC Homo sapiens cDNA clone GKCDRE12 5'
1933	11828	21710		3.0E-48	4885170 NT	L	Homo sapiens chromoscme X open reading frame 6 (CXORF6) mRNA
1933	11828		18.97	3.0E-48	4885170 NT	LN	Homo sapiens chromosome X open reading frame 6 (CXORF6) mRNA

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Single |

hi14b12.x1 NCI_CGAP_GU1 Homo sapiens cDNA clone IMAGE:2972255 3' similar to SW:DCRB_HUMAN nv03f05.s1 NCI_CGAP_Pr22 Homo sapiens cIDNA clone IMAGE:1219137 3' similar to contains PTR5.b1 Homo sapiens amyloid beta (A4) precursor profein (protease nexin-II, Alzheimer disease) (APP), mRNA CBAP1D3842 Pediatric pre-B cell acute lymphoblastic leukemia Baylor-HGSC project=TCBA Homo Homo sapiens v-rel avian reticuloendotheliosis viral oncogene homolog A (nuclear factor of kappa light UI-H-BW1-ant-a-10-0-UI st NCI_CGAP_Sub7/Homo sapiens cDNA clone IMAGE:3082267 3* fmfc7 Regional genomic DNA specific cDNA library Homo sapiens cDNA clone CR17-26 2x80c03.11 Soares ovary tumor NbHOT Homo sapiens cDNA clone IMAGE:810052 5' 601305064F1 NIH_MGC_39 Homo sapiens cDNA clone IMAGE:3639782 5' Homo sapiens cisplatin resistance-associated overexpressed protein (LOC51747), mRNA no18g01.s1 NCI_CGAP_Phe1 Homo sepiens ¿DNA clone IMAGE:1101072 3 no18g01.s1 NCI_CGAP_Phe1 Homo sepiens ¿DNA clone IMAGE:1101072 3 601888096F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:4122119 6' Homo sapiens dopamine transporter (SLCBA3) gene, complete cds Homo sapiens dopamine transporter (SLCBA3) gene, complete cds 15d6 Human retina cDNA randomly primed subilbrary Homo sapiens cDNA Human endogenous retroviral DNA (4-1), complete retroviral segment P56555 DOWN SYNDROME CRITICAL REGION PROTEIN B. Top Hit Descriptor polypeptide gene enhancer in B-celis 3 (p65)) (RELA), mRNA MR4-BT0657-060400-201-e10 BT0657 Homo saplens cDNA Homo sapiens RNA binding motif protein 6 (REM6) mRNA Homo sapiens chromosome 21 segment HS21,0102 Homo sapiens huntingtin (Huntington disease) (HD) mRNA Homo sapiens mRNA for KIAA1501 protein, partial cds Homo saplens mRNA for KIAA1245 protein, pertial cds Homo sapiens mRNA for KIAA1501 protein, partial cds AV743451 CB Homo sapiens cDNA clone CBCCGG1 Homo saplens chromosome 21 segment HS21,C046 Homo sapiens B cell linker protein (SLP65), mRNA Homo sapiens B cell linker protein (SLP65), mRNA Mus musculus T-box 20 (Tbx20), mRNA sapiens cDNA clone TCBAP3842 PTR5 repetitive element EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN HUMAN EST_HUMAN EST_HUMAN EST HUMAN HUMAN HUMAN HUMAN Top Hit Database Source EST EST 눌 뉟 11496238 NT 7708534 NT z 눋 4602166|NT 5032032|NT z 10048417|NT 11429808 Top Hit Acession 1.0E-48 AF119117.1 1.0E-48 W26785.1 8.0E-49 10048 1.0E-48 AB033071.1 1.0E-48 BF304683.1 AW664531.1 2.0E-48 AA613171.1 2.0E-48 AA613171.1 2.0E-48 1149 2.0E-48 AV743451.1 2.0E-48 AA465007.1 2.0E-48 AB040934.1 1.0E-48 AL163302.2 1.0E-48 AL163246.2 BE084571.1 3.0E-48 AA659930.1 3.0E-48 BF514170.1 2.0E-48 AA631940.1 2.0E-48 BE246065.1 BE737164.1 AF119117.1 ġ 1.0E-48 M10976.1 1.0E-48 .0E-48 / 3.0E-48 3.0E-48 2.0E-48 1.0E-48 2.0E-48 1.0E-48 1.0E-48 1.0E-48 1.0E-48 (Top) Hit BLAST E dost Simila 2.39 0.88 2.35 2.86 72.29 4.29 2.45 1.63 1.23 4.48 5.06 5.08 2.95 1.18 1.35 6.93 2.21 5.72 1.56 0.85 3.26 Expression Signal 28376 25539 25540 26325 27586 28023 23284 24114 28464 26473 24828 25070 27404 25702 19827 19844 21032 ORF SEQ 26465 ÖΝΘ 16743 16308 13483 16768 18128 14328 15082 19579 16167 15507 10027 15469 15469 16301 14952 18802 19606 15601 SEQ ID 10037 1630 ÿ Probe SEQ ID 6889 8248 5553 5553 9184 3443 5082 6303 7526 7934 8999 8999 9145 5692 3579 5592 6440 6447 6864 822 8 4431 ස 1274 1876 7337 8440 9511 ÿ

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Top Hit Descriptor	Mus musculus T-box 20 (Tbx20), mRNA	Human inositol 1,4,5 trisphosphate receptor type 1 mRNA, partial cds	Homo sapiens proteasome (prosome, macropain) 26S subunit, ATPase, 4 (PSMC4) mRNA	Homo sapiens proteasome (prosome, macropain) 26S subunit, ATPase, 4 (PSMC4) mRNA	Homo saplens proteasome (prosome, macropain) 26S subunit, ATPase, 4 (PSMC4) mRNA	Homo sapiens proteasome (prosome, macropain) 26S subunit, ATPase, 4 (PSMC4) mRNA	Homo saplens proteasome (prosome, macropain) 26S subunit, ATPase, 4 (PSMC4) mRNA	Homo saptens proteasome (prosome, macropain) 26S subunit, ATPase, 4 (PSMC4) mRNA	Homo sapiens chromosome 21 segment HS21:0084	HYPOTHETICAL PROTEIN DJ845024.3	wf25h04.x1 Soares_NFL_T_GBC_S1 Homo seplens cDNA clone IMAGE:2359683 3' similar to TR:O54923	VO14825 TOECTS.	DKFZp762C033_s1 762 (synonym: hmel2) Homo sepiens cDNA clone DKFZp762C033 3'	ba55g05.x1 NIH_MGC_10 Homo saptens cDNA clone IMAGE:2900504.3' similar to gb:X17208.40S RIROSOMAI PROTFIN S4 (HI MAN): ch:M20632 Mouse I I Ren3 protein mRNA from a renetitive element	complete (MOUSE);	UI-H-Bi3-alo-a-05-0-UI.s1 NCI_CGAP_Sub5 Homo sapiens cDNA clone IMAGE:3068048 3'	EST77525 Pancreas tumor III Homo sapiens cDNA 5' end	EST77525 Pancreas tumor III Homo sapiens cDNA 5' end	zj29c08.s1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clane IMAGE:451694 3'	Homo sapiens chromosome 21 segment HS21C010	Homo sapiens chromosome 21 segment HS21,C010	zp29c07.r1 Stratagene neuroepithelium (#937231) Homo sapiens cDNA clone IMAGE:610860 5' similar to ; TR:G233226 G233226 RTVL-H PROTEIN ; contains LTR7.t3 LTR7 trepetitive element ;	Homo sapiens putative tumor suppressor ST13 (ST13) mRNA, complete cds	Homo sapiens similar to ribosomal protein S27/(metallopanstimulin 1) (H. sapiens) (LOC63362), mRNA	x08b01.x1 NCI_CGAP_Ut4 Homo saplens cDNA clone IMAGE:2675593 3' similar to WP:B0350.2B	CECOLOS	zr90f05.r1 NCL_CGAP_GCB1 Homo sapiens cDNA clane IMAGE:682977 5	Homo sapiens glutathione S-transferase theta 2 (GSTT2) and glutathione S-transferase theta 1 (GSTT1) genes, complete cds	H.sapiens mRNA for acetyl-CoA carboxylase
Top Hit Database Source	۲	TN	NT	IN.	۲	<u>F</u>	M	IN	FX	SWISSPROT	HAMILI FOR	NAWOU CO	EST_HUMAN		EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	۲	LN	EST_HUMAN	님	FZ	111111111111111111111111111111111111111	ES HOMAN	EST_HUMAN	TN	NT
Top Hit Acessian No.	10048417 NT	U23850.1	5729990 NT	AL163284.2	060811	4 404 4		AL120937.1		AW731740.1	AW 452218.1	AA366556.1	AA366556.1	AA707567.1	AL163210.2	AL163210.2	AA172121.1	U17714.1	11436355 NT		AW189533.1	AA210798.1		X68968.1					
Most Similar (Top) Hit BLAST E Value	8.0E-49	8.0E-49	7.0E-49	7.0E-49	7.0E-49	7.0E-49	7.0E-49		7.0E-49	7.0E-49	100	87-JO:/	7.0E-49		6.0E-49	6.0E-49	6.0E-49	8.0E-49	6.0E-49	5.0E-49	5.0E-49	5.0E-49	6.0E-49	5.0E-49		ı	4.0E-49	4.0E-49	3.0E-49
Expression Signal	2:35	3.19	1.47	1.47	1.74	1.74	1.99	1.99	4.05	0.95	8	3	1.34		11.77	2.92	2.69	2.69	3.43	3.37	3.37	194	5.18	5.13	0, 10	37.45	2.43	3.3	0.93
ORF SEQ ID NO:	25703	26900	20157	20158	20157	20158	20157		20958	24227	. 00,90		26107		19979		29022	29023		20452	20453	21524	22477	22957		20208			20298
Exan SEQ ID NO:	15601	16706	10335	10335	10335	10335	10335	10335	11112	14443	45074	1/701	16277		10162	18467	18728	18728	19498	10628	10628	11653	12583	13159	l _		19710		10489
Probe SEQ ID NO:	5692	6827	134	134	388	388	389	389	1202	4650	5254	2	5357		190	8600	8920	8920	9507	695	969	1753	2721	3235		314	9376	9469	548

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Most Similar (Top) Hit Top Hit Acession Database Source	1.1 EST_HUMAN	2.08 3.0E-49 U46999.1 NT Human type IV collagen (COL4A6) gene, exon 40	9.6 3.0E-49 H39479.1 EST_HUMAN EST25e12 WATM1 Homo saplens cDNA clone 25e12		Г	1.4 2.0E-49 N28446.1 EST_HUMAN yx23d06.11 Soares melanocyte 2NbHM Homo explens cDNA clone IMAGE:282571 5		oz88402.x1 Soares_senescent_fibroblasts_NbHSF Homo sapiens cDNA clone IMAGE:1682403 3' similar to gb:M31470 RAS-LIKE PROTEIN TC10 (HUMAN);contains element;contains element MER22	1.12 2.0E-49 A1167357.1 EST_HUMAN repetitive element;	BF511846.1 EST_HUMAN	AV717838.1 EST_HUMAN	1.97 2.0E-49 M86033.1 EST_HUMAN EST02558 Fetal brain, Strategene (cat#936209) Homo sepiens cDNA clone HFBCY50	AF163864.1 NT	3.95 1.0E-49 BF03527.1 EST_HUMAN 601458531F1 NIH_MGC_68 Homo sapiens cDNA clone IMAGE:3862086 5'		1.0E-49 BF131007.1 EST_HUMAN		1.0E-49 BE398110.1 EST_HUMAN	1.0E-49 N25884.1	1 OF 40 N2RB84 1	1.0E-49 11321580 NT	1.28 1.0E-49 11321580 NT Homo sepiens succinate-CoA ligase, GDP-forming, eipha subunit (SUCLG1), mRNA	T HUMAN	1.0E-49 AL043129.2	1.0E-49 11427366	1.73 1.0E-49 BE159343.1 EST_HUMAN MRO-HT0407-010200-006-f02 HT0407 Homo/saplens cDNA	11418322	4F101475.1 NT	4L163202.2 NT	1.89 8.0E-50 X95097.2 NT Homo saplens mRNA for VIP receptor 2
Most Similar (Top) Hit BLAST E Vaiue									2.0E-49		İ	L							1.0E-49	4 OF 40				L						
Expression Signal					2.66							1.97	1.53	3.95	2.58	4.97										1.73	2	1.08	2.59	
ORF SEQ ID NO:		24584				22914			24373	L					21531	25011	26307												19951	
Exan SEQ ID NO:	12479	14788	16248	18487	10582	13110	L		14579	14590	15943	16597	L	10807	11659	15210	16151	L	L .			Ŀ	L		L		<u> </u>	14802	10136	L
Probe SEQ ID NO:	2611	4909	6386	8621	645	3185	3521		4693	4704	6040	6717	9467	881	1760	5289	6287	6287	6322	8277	6709	6209	7200	7830	8630	9018	9367	4923	163	702

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Single Extri Flobes Expressed in Lean	Top Hit Descriptor	Homo sapiens mRNA for VIP receptor 2	Homo sapiens homogentisate 1,2-dloxygenase gene, complete cds	Homo sapiens actinin, alpha 1 (ACTN1) mRNA	Homo sapiens p47 (LOC51674), mRNA	Homo sapiens p47 (LOC51674), mRNA	Homo sapiens capping protein (actin filament) inuscle Z-line, beta (CAPZB), mRNA	np82d06.s1 NCI_CGAP_Br2 Homo sepiens cDNA clone IMAGE:1130891 3' similar to gb:J05459 GLUTATHIONE S-TRANSFERASE TESTIS/BRAIN (HUMAN);	QV0-BT0703-280400-211-e08 BT0703 Homo sepiens cDNA	wm55g11.x1 NCI_CGAP_Ut2 Homo saplens cDNA clone IMAGE:2439908 3'	ho36h04x1 NCL_CGAP_Ut1 Homo sapiens cDNA done IMAGE:3039511 3' similar to contains MER29.b3 MFR29 repetitive element :	EST182775 Jurkat T-cells VI Homo sapiens clONA 5' end	EST182775 Jurkat T-cells VI Homo saplens cDNA 5' end	CM0-BT0792-300500-398-b05 BT0792 Homo saplens cDNA	CM0-BT0792-300500-398-b05 BT0792 Homo sapiens cDNA	nl45h10.s1 NCI_CGAP_Pr4 Homo sapiens cDNA clone IMAGE:1043683 similar to contains PTR5.t3 PTR5	repetitive element;	zt62b01,r1 Soares_testis_NHT Homo sapiensicDNA clone IMAGE:726889 5' similar to TR:G1335769 G1335769 GAG-POL POLYPROTEIN. ;	no54e09.s1 NCI_CGAP_SS1 Home saplens cDNA clone IMAGE:1104520 3' similar to gb:X53741_ma1 FIRLII NJ-1 ISOFORM A PRECURSOR (HUMAN):	Human endogenous retrovirus RTVL-H2	obo3f08.s1 NCI_CGAP_Kid3 Homo sapiens cDNA clone IMAGE:1322627 3'	CMYA5 Human cardiac muscle expression library Homo sapiens cDNA clone 4151935 similar to CMYA5	Cardiomyopathy associated gene 5	Homo sepiens similar to sema domain; immunoglobulin domain (Ig), short basic domain, secreted,	(semaphorin) 3A (H. sapiens) (LOC83232), mKNA	Homo sapiens FYVE domain-containing dual specificity protein phosphatase FYVE-DSP1a mRNA, complete	spo	Homo sapiens FYVE domain-containing dual specificity protein phosphatase FYVE-DSP1a mRNA, complete	spo	Homo sapiens mRNA for KIAA1598 protein, partal cds	Homo sapiens CTL2 gene
ופ באמוו גוממפא	Top Hit Database Source	H	H IN					EST_HUMAN G	EST_HUMAN Q	EST_HUMAN W	M NAMI IL TOTA	T	HUMAN	HUMAN	EST HUMAN C		EST_HUMAN re	EST HUMAN G		Т	T HUMAN	Т	EST_HUMAN C			Ξ_	Ā				H
Dillo Dillo	Top Hit Acession No.	95097.2	8.0E-50 AF000573.1	4501890 NT	7706394 NT	7706394 NT	4826658 NT	8.0E-50 AA633467.1					Γ	-332038.1	-332938.1		5.0E-50 AA557883.1	5 0E-50 AA403053.1		T	3.0E-50 AA746142.1	Γ	3.0E-50 AW755254.1		11421514 NT		3.0E-50 AF233436.2		F233436.2	8046818.1	J245621.1
	Most Similar (Top) Hit BLAST E Value	8.0E-50 X	8.0E-50	8.0E-50	8.0E-50	8.0E-50	8.0E-50	8.0E-50	7.0E-50	7.0E-50 /	20 20 20	8.0E-50 /	8.0E-50			l	5.0E-50	5 05-50	7 20 70	3 05-50	3.0E-50				3.0E-50		3.0E-50			3.0E-50 A	
	Expression Signal	1.89	121	2.51	1.36	1.36	1.69	2.1	0.96	9.52	27.7	3.17	3.17	0.85	0.85		5.64	1.85	,	27.45	1.14		4.6		1.55		4.01		4.01		5.94
	ORF SEQ ID NO:	20461		21497	١.	22205		28811				28326		Ĺ	21523			29088	1		22981		23392		26180		26540			27721	
	Exan SEQ ID NO:	10635	10934	11628	L	ł	ł		L	Ł		L	<u></u>	L	L	Ŀ	17132	18795	1.	L		1_	13606	ł	16039		16364	1	16364	17499	17909
	Probe SEQ ID NO:	702	1016	1727	2432	2432	2666	8711	603	8139	2070	8180	8190	1752	1752		7255	Raga	6	4808	3259		3692		9909		6505		6505	7649	8760

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Top Hit Descriptór	C dass 1 region	Homo sapiens midline 1 (Opitz/BBB syndrome) (MID1) mRNA	Homo sapiens decorin D mRNA, complete cds, alternatively spliced	Human HALPHA44 gene for alpha-tubulin, exons 1-3	Human HALPHA44 gene for alpha-tubulin, exons 1-3	Mus musculus keratin complex 2, gene 6g (Krt2-6g), mRNA	Mus musculus keratin complex 2, gene 6g (Krt2-6g), mRNA	PM3-BN0137-290300-002-g11 BN0137 Homo sepiens cDNA	PM3-BN0137-290300-002-q11 BN0137 Homo sapiens cDNA	Homo sapiens chromosome 21 segment HS21C009	Homo sapiens Xq pseudoautosomal region; segment 1/2	zk51c09.r1 Soares_pregnant_uterus_NbHPU l/lomo sapiens cDNA clone IMAGE:486352 5	np88e09.s1 NCI_CGAP_Lu1 Homo sapiens cDNA clone IMAGE:1142440 3' similar to gb:X12671_ma1 HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN A1 (HUMAN);	Homo sapiens PDZ-73 protein (PDZ-73/NY-CQ-38), mRNA	AU138590 PLACE1 Homo saplens cDNA clone PLACE1008887 5'	QV4-NT0028-200400-180-d05 NT0028 Homo sapiens cDNA	xn34a03.x1 NCI_CGAP_Kid11 Homo sapiens 5DNA clone IMAGE:2695564.3' similar to TR:Q9Z340 Q9Z340 ATYPICAL PKC SPECIFIC BINDING PROTEIN ;	DKFZp434B2229_r1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434B2229 5'	DKFZp434B2229_r1 434 (synonym: htss3) Homo sapiens cDNA clone DKFZp434B2229 5'	UI-H-BW0-alp-b-05-0-UI.s1 NOI_CGAP_Sub6 Homo saplens cDNA clone IMAGE:2729817 3'	Homo sepiens KIAA0929 protein Msx2 Interacting nuclear target (MINT) homolog (KIAA0929), mRNA	Homo sapiens KIAA0929 protein Msx2 interacting nuclear target (MINT) homolog (KIAA0929), mRNA	Homo sapiens solute carrier family 2 (facilitated glucose transporter), member 9 (SLC2A9), mRNA	Homo sapiens solute carrier family 2 (facilitated glucose transporter), member 9 (SLC2A9), mRNA	n related (Hpr) gene exon 3	Homo sapiens milogen-activated protein kinase kinase 1 (MKK4) gene, exon 4	Homo sapiens mitogen-activated protein kinase kinase 1 (MKK4) gene, exon 4	Homo sapiens cerebral cell adhesion molecule((LOC51148), mRNA	orotein (B9), mRNA	Homo sapiens interleukin 17 receptor (IL17R), mRNA
	Homo sapiens MHC class 1 region	sapiens midline 1 (Opitz/BBB syr	sapiens decorin D mRNA, comple	n HALPHA44 gene for alpha-tubu	n HALPHA44 gene for alpha-tubu	nusculus keratin complex 2, gene	nusculus keratin complex 2, gene	3N0137-290300-002-911 BN0137	3N0137-290300-002-911 BN0137	sapiens chromosome 21 segmen	sapiens Xq pseudoautosomal reg	09.r1 Soares_pregnant_uterus_N	09.s1 NCI_CGAP_Lu1 Homo sap ROGENEOUS NUCLEAR RIBOI	sapiens PDZ-73 protein (PDZ-73	8590 PLACE1 Homo sapiens cDN	VT0028-200400-180-d05 NT0028	03.x1 NCI_CGAP_Kid11 Homo s 40 ATYPICAL PKC SPECIFIC BI	p434B2229_r1 434 (synonym: hte	p434B2229_r1 434 (synonym: hte	3W0-alp-b-05-0-UI.s1 NCI_CGAF	sepiens KIAA0929 protein Msx2	sapiens KIAA0929 protein Msx2	sapiens solute carrier family 2 (fa	sapiens solute carrier family 2 (fa	Human haptoglobin related (Hpr) gene exon 3	sapiens mitogen-activated protein	sapiens mitogen-activated protein	sapiens cerebral cell adhesion mo	Homo sapiens B9 protein (B9), mRNA	sapiens interleukin 17 receptor (il
Top Hit Database Source								HUMAN	HUMAN			EST_HUMAN 2k51c	HUMAN		T HUMAN	EST_HUMAN QV4-	EST_HUMAN Q9Z3	_		EST_HUMAN UI-H-										
Top Hit Acession No.	AF055066.1 NT	4557752 NT	NF138303.1 NT	K06956.1 NT	X06956.1	9910293 NT	9910293 NT	3E007080.1 EST	3E007080.1 EST	4L163209.2 NT	4J271735.1 NT	4A043738.1 ES	4A610842.1 EST	11439587 NT	4U138590.1 ES	4W889219.1 ES	4W274720.1 ES	4L079628.1 ES	4L079628.1 ES	4W295603.1 ES	7657266 NT	7657266 NT	9910553 NT	9910553 NT	X01788.1 NT	AF070083.1 NT	4F070083.1 NT	11429665 NT	7661535 NT	11526289 NT
Most Similar (Top) Hit BLAST E Value	1 -	2.0E-50		2.0E-50 X		2.0E-50	2.0E-50	1.0E-50 B	1.0E-50 B	_	1.0E-50 A	9.0E-51 A		8.0E-51		7.0E-51 A	7.0E-51 A	7.0E-51	7.0E-51 A	7.0E-51 A	6.0E-51	6.0E-51	6.0E-51	6.0E-51	6.0E-51	6.0E-51 A	6.0E-51 A	6.0E-51	6.0E-51	6.0E-51
Expression Signal	16:4	4.6	18.02	6.27	6.27	1.53	1.63	+	4	2.1	8.98	1.22	4.89	2.34	1.28	1.36	0.83	1.26	1.26	2.38	5.3	12.92	0.78	0.78	2.26	69.6	6.68	2.16	2.26	1.72
ORF SEQ ID NO:		20823	21197	26995	26938		27762	20018	20019			27450	24151			22967	23042	23757		23927	21714	23150	23887		25651	25656	25657	24859		28716
Exan SEQ ID NO:	10691	10979	11330	16802	16802		17538	10204	10204	10398	12195	17244	14361	16369	L	L	13238		13978	14153	11831	13345	14110	<u>L</u>	15558	15562	15562	15116		18448
Probe SEQ ID NO:	761	1063	1425	6924	6924	7686	7686	235	235	454	2314	7375	4467	6510	7448	3246	3317	4078	4076	4254	1936	3428	4212	4212	5645	5850	5650	6168	7583	8580

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Most Similar Top Hit Acession (Top) Hit Top Hit Acession Signal BLASTE No. Signal Value	10.92 6.0E-61 AL163203.2 NT	1.47 5.0E-51 4507500 NT	1.37 6.0E-51 AL133204.1 NT	0.84 6.0E-61 6031980 NT	22311 11.48 5.0E-51 AJ007559.1 NT Homo sepiens mRNA for nucleoparin 155	23568 1.08 6.0E-51 M30838.1 NT Human Ku (p70/p80) subunit mRNA, complete cds	1.08	.1 NT	28739 3.72 6.0E-51 5803138 NT Homo sepiens RNA binding motif protein 3 (RB/M3), mRNA	19926) 0.92 3.0E-51 A1587348.1 EST HUMAN KERATIN, TYPE I CYTOSKELETAL 18 (HUMAN);	4.16 3.0E-51 AI587348.1	Į.	1.73	3.0E-51 M29063.1	1.58 3.0E-51 AF003528.1 NT regions - 1.58 3.0E-5	Homo saplens ubiquitin protein ligase E3A (human papilloma virus E6-associated protein, Angelman syndrome) (UBE3A) mRNA syndrome) (UBE3A) mRNA	1.08 2.0E-51 BE391063.1 EST_HUMAN	20425 1.08 2.0E-51 BE391063.1 EST HUMAN 601285694F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3607463 5'	2.24 2.0E-51 AA233352.1 EST_HUMAN	2.21 2.0E-61 AI492416.1 EST_HUMAN	1.02 2.0E-51 AW137828.1	EST_HUMAN	1.61 2.0E-61 BE901994.1 EST_HUMAN	EST_HUMAN	1.88 2.0E-51 AI917078.1 EST_HUMAN	27600 5.25 2.0E-51 BE165980.1 EST HUMAN MR3-HT0487-150200-113-g01 HT0487 Homo saplens cDNA
ORF SEQ Express ID NO: Signa						23566	23567	24643	28739	19926	20917	23906	28500			20139	20424	20425	21431	23373	24071	25670	27114	27115	27662	27600
SEQ ID	Ш	786 10715			2548 12422	3863 13774		5004 14879	8601 18468	130 10104		4232 14130	<u> </u>	7131 17008	9675 19233	362 10318	673 10607	673 10607	1	3672 .13586	4392 14288	L	7047 16924	7047 16924		7539 17390
Probe SEQ ID NO:						ຼິ		အ	8					_						Ľ	4	Ľ	Ľ	Ľ		

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Table 4
Single Exon Probes Expressed in Heart

	Т	Г				Г	_						Т	Т	Т	7	П			Τ"	T	Γ		7,10	Ï	Πĺ				7	_	T
Top Hit Descriptor	H.eapiens flow-sorted chromosome 6 HindIII fragment, SC6pA18H7	Homo sapiens SH3-containing protein SH3GLB1 mRNA, complete cds	Homo sapiens nucleoporin 155kD (NUP155) mRNA	Homo sapiens T-cell lymphoma invasion and metastasis 1 (TIAM1) mRNA	601440687F1 NIH_MGC_72 Homo saplens cDNA clone IMAGE:3915836 5'	Hömo sapiens hydroxystercid (17-beta) dehydrogenase 4 (HSD17B4), mRNA	Homo sapiens Ran GTPase activating protein 1 (RANGAP1), mRNA	Homo sapiens DNA for Human P2XM, complete cds	Homo sapiens hypothetical protein FLJ10675 (FLJ10675), mRNA	Human endogenous retroviral DNA (4-1), complete retroviral segment	Human endogenous retroviral DNA (4-1), complete retroviral segment	bb66b07.y1 NIH_MGC_9 Homo saplens cDNA clone IMAGE:3030421 5' similar to gb:X16493 M.musculus mbNA for 2nf 1 min finds mortal AAO ISEN	FINANCE DE SPIRIT MICH MICH AND SERIORS FINANCE MARCHASASA FI	Mound human again manning to chromosome 20. similar to membrane transporters	Nova nama garo napping to circumsome ze, simila to maina anaspecta s	II.3-C10214-231299-053-E12 CT0214 Homo sapiens cDNA	Homo sapiens interleukin 21 receptor (IL21R), mRNA	Macaca mulatta beta-tubulin mRNA, complete cds	Homo saplens NADH dehydrogenase (ubiquinche) Fe-S protein 5 (15kD) (NADH-coenzyme Q reductase) (NDLIESS) mRNA	Homo sabiens SET domain and mariner transposase fusion gene (SETMAR) mRNA	Homo sapiens SET domain and mariner transposase fusion gene (SETMAR) mRNA	wj49c04.x1 NCI_CGAP_Lu19 Homo sapiens cDNA clone IMAGE:2406150 3' similar to contains THR.b2	THR repetitive element;	wj49c04.x1 NOI_CGAP_Lu19 Homo sapiens cDNA clone IMAGE:2406150 3' similar to contains THR.b2	THR repetitive element ;	AV715377 DCB Homo saplens cDNA clone DCBAIE03 5'	zd49g12.r1 Soares_fetal_heart_NbHH19W Homo sapiens cDNA clone IMAGE:344038 5'	Homo sapiens LIM domain kinase 2 (LIMK2), niRNA	xn72e07.x1 NCI_CGAP_CML1 Homo sapiens cDNA clone IMAGE:2700036 3' similar to contains Alu	repetitive element;contains element LTR2 repetitive element ;	wf67d05x1 Soares_NFL_T_GBC_S1 Homo sipiens cDNA clone IMAGE:2360649 3' similar to TR:Q16859 Q16859 CARBOXYLESTERASE;	zu75h12.s1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:743879 3'
Top Hit Database Source	¥	LN	FZ	Į.	EST_HUMAN	N	N	Z L	LN	FZ	N	TOTAL TOTAL	EST TIMAN	LO LICENSIA	- II	EST_HUMAN	NT	LN		L L	Z		EST_HUMAN		EST_HUMAN	EST_HUMAN	EST_HUMAN	FZ		EST_HUMAN	EST HUMAN	EST_HUMAN
Top Hit Acession No.	Z78898.1	AF257318.1	4758843 NT	4507500 NT	BE622032.1	11417035 NT	11418177 NT	AB002059.1	11437042 NT	M10976.1	M10976.1	7 242404	DE20/0/5.1	A1 197100 3	4L13/100.3	AW848041.1	11141868 NT	AF147880.1	TN 087826A	5730038 NT	5730038 NT		A1831462.1		AI831462.1	AV715377.1	W70260.1	11417990 NT		AW 236297.1	Al808985.1	AA634445.1
Most Similar (Top) Hit BLAST E Value	5.0E-52	4.0E-52	4.0E-52	4.0E-52	4.0E-52	4.0E-52	4.0E-52	4.0E-52	3.0E-52		2.0E-52		2.0E-52			2.0E-62	2.0E-52	2.0E-52	2.50	2 OF-52	2.0E-52		2.0E-52		2.0E-52	2.0E-52	2.0E-52	2.0E-52		2.0E-52	2.0E-52	
Expression Signal	2.07	0.93	8.58	0.82	1.24	7.25	4.25	5.09	86.6	1.39	1.39	1	0/:	3.52	7/7	2.74	1.49	8.39	60	4 53	4.53		5.33		5.33	3.09	2.08	3.22		8.86	4.28	1.37
ORF SEQ ID NO:	24021			L						20299	20300	0000	07777				25860				27919		28671			28689				24894		20276
Exan SEQ ID NO:	14237		11648	13761	16566	16843	18992	19293	13908	10491	10491	}	1	1	14/07	15416	15747	16905	17087	1	1		18408	J	18408	18419	18548	18701		19755	19112	
Probe SEQ ID NO:	4340	1639	1748	3850	9899	6965	9291	8778	4002	550	550	276	02420	2007	480%	5497	5841	7028	7458	7825	7825		8536		8536	8547	8659	889-1		9101	9496	522

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	Top Hit Descriptor	Homo saplens giutamate-ammonia ligase (glutamine synthase) (GLUL) mRNA	Homo sapiens aryisulfatase D (ARSD), transcript variant 1, mRNA	polereverse transcriptase homolog (retroviral element) [human, endogenous retroviral element RTVL-Hp1,	Genomic, 660 nt]	Human P-glycoprotein (MDR1) gene, exon 4	Human PMS2 related (hPMSR2) gene, complete cds	Human aldolase C gene for fructose-1,6-bisphosphate aldolase	Homo sapiens chromosome 21 segment HS21C027	Homo capians chromosome 21 segment HS210002	Homo sapiens protein tyrosine phosphatase PTPCAAX1 (hPTPCAAX1) mRNA, complete cds	Homo sapiens proteasome (prosome, macropain) subunit, beta type, 2 (PSMB2), mRNA	Homo sapiens protein kinase, cAMP-dependent, regulatory, type II, beta (PRKAR2B) mRNA	Homo sapiens core binding factor alpha1 subunit (CBFA1) gene, exon 3	Homo sapiens predicted osteoblast protein (GS3788), mRNA	601804771F1 NIH_MGC_54 Homo sapiens cDNA clone IMAGE:413Z793 5'	ば4f07x1 NCI_CGAP_Brn23 Homo sapiens cDNA clone IMAGE:2099077 3' similar to contains THR.t1	THR repetitive element ;	Homo sapiens heterogeneous nuclear ribonucleoprotein C (C1/C2) (HNRPC) mRNA	RC3-ST0197-151099-011-g10 ST0197 Homo saplens cDNA	Homo sapiens chromosome 21 segment HS21C085	Homo sapiens chromosome 21 segment HS21C085	Homo sapiens hook1 protein (HOOK1), mRNA	601810969F1 NIH_MGC_48 Homo sapiens cDNA clone IMAGE:4053977 5'	601810969F1 NIH_MGC_48 Hamo sapiens clONA clone IMAGE:4053977 5'	Homo sapiens DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes,	complete cds)	wz22c07.x1 Soares_Dteckgreafe_colon_NHCD Homo sapiens cDNA clone IMAGE:2558796 3'	IL2-UM0081-240300-055-D03 UM0081 Homo sapiens cDNA	QV3-BT0381-270100-073-d06 BT0381 Homo saplens cDNA	GIF=growth Inhibitory factor [human, brain, Génomic, 2015 nt]	Homo sapiens FGFR1 oncogene partner (FOP), mRNA	EST77525 Pancreas fumor III Homo sapiens ¢DNA 5' end	Homo sapiens Bruton's tyrosine kinase (BTK)∖ alpha-D-galactosidase A (GLA), L44-like ribosomal protein (L44L) and FTP3 (FTP3) genes, complete cdṡ	
160 H	Database Source	N-	LZ LZ		NT	LN L	TN	뒫	Į.	NT	NT	NT	Z	٦	N	EST_HUMAN		EST_HUMAN	NT	EST_HUMAN	M	NT	TN	EST_HUMAN	EST_HUMAN		L	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	IN	EST_HUMAN	Į.	
	Top Hit Acession No.	4504026 NT	4502238 NT		S61070.1	M29426.1	U38964.1	X07292.1	AL163227.2	AL163202.2	U48296.1	11426321 NT	4506084 NT	AF001446.1	7861713 NT	BF238465.1		AI421782.1	4758543 NT	AW813563.1	AL163285.2	AL163285.2	7705414 NT	BF128701.1	BF128701.1		AB026898.1	AW050836.1	AW803563.1	BE069344.1	S72043.1	5901953 NT	AA366556.1	U78027.1	
-	(Top) Hit BLAST E Vafue	1.0E-52	1.0E-62				1.0E-52	1.0E-52	1.0E-52	1.0E-52	1.0E-52	1.0E-52	9.0E-53	9.0E-53	9.0E-53	7.0E-53		7.0E-53	5.0E-53	5.0E-53	4.0E-53	4.0E-53	4.0E-53	4.0E-53	4.0E-53		3.0E-53	3.0E-53	3.0E-53	3.0E-53	3.0E-53	3.0E-53		2.0E-53	
	Expression Signal	8.25	1.2		1.41	3.69	2.11	3.18	1.64	1.61	1.84	2.04	1.03	1.01	0.93	2.08		2.98	2.2	1.72	1.92	1.92	1.09	3.33	3.33		2.59	1.29	1.18	0.85	98.8	8.59	4.25		
	ORF SEQ ID NO:	21111	_		22741	24967			l		28283		23420		24611				23690		19834	19835	24387	28685	28686									22060	
Exe	SEQ ID	11255	l	l	12949	15192	15757	16256	16810	17873	18035	18094	13635	14191	l_	19018		19632	13915	19048	10031	10031	l	18417	18417		12493	13584		L	16712	L	ı		ı
Probe	SEQ ID	1349	2489		3021	5270	5851	6394	6932	8023	8147	8210	3723	4293	4966	9338		9752	4009	9389	£	43	4715	8645	8545		2825	3670	4486	4833	8833	7238	450	2279	

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							,	_				, .				-		4	,,,		سا: ,	مدع					- 4.		1-41	41-49 41-49
Top Hit Descriptor	Homo sapiens ATPase, H+ transporting, lysosomal (vacuolar proton pump) 31kD; Vacuolar proton-ATPase, subunit E; V-ATPase, subunit E (ATP6E), mRNA	Homo sapiens core-binding factor, runt domain, alpha subunit 2; translocated to, 1; cyclin D-related (CBFAZT1) mRNA	Homo saplens core-binding factor, runt domain, alpha subunit 2; translocated to, 1; cyclin D-related (CBFA271) mRNA	Homo sapiens dihydropyridine receptor alpha 2 subunit (CACNA2D1) gene, exon 6	Human Krueppel-related DNA-binding protein (IFS4) gene, partial cds	Homo sapiens SKAP55 homologue (SKAP-HOM) mRNA	Homo sapiens chromosome 21 segment HS21 C081	Homo sapiens chromosome 21 segment HS210081	PM1-CT0396-170800-001-g03 CT0396 Homo sapiens cDNA	PM1-CT0398-170800-001-g03 CT0396 Homo saplens cDNA	2822665.5prime NIH_MGC_7 Homo sapiens cDNA clone IMAGE:2822665 5'	Homo saplens Xq pseudoautosomal region; segment 2/2	Homo saplans DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes,	complete cas)	CM4-NN1029-150800-543-e02 NN1029 Homo saptens cDNA	H.sapiens mRNA for hnRNPcore protein A1	Homo sapiens T-cell lymphoma invasion and metastasis 1 (TIAM1) mRNA	Homo sapiens T-cell lymphoma invasion and metastasis 1 (TIAM1) mRNA	Homo sapiens IQ motif containing GTPase activating protein 1 (IQGAP1) mRNA	601272863F1 NIH_MGC_20 Homo saplens cDNA clone IMAGE:3614031 5'	Homo sapiens insulin-like growth factor 2 receptor (IGF2R) mRNA	Homo sapiens ubiquitin specific protease 13 (isopeptidase T-3) (USP13) mRNA	Homo saplens ubiquitin specific protease 13 (isopeptidase T-3) (USP13) mRNA	Homo sapiens ubiquitin specific protease 13 (isopeptidase T-3) (USP13) mRNA	Homo sapiens ublquitin specific protease 13 (isopeptidase T-3) (USP13) mRNA	Homo saplens ATP-binding cassette, sub-family A (ABC1), member 8 (ABCA8), mRNA	ai76c12.s1 Soares_testis_NHT Homo sapiens cDNA clone 1377046 3' similar to contains MER30.t3 MER30	Jepannya digitali,	Homo sapiens mknA for monocyte chemotacti⇒ protein-2	yw68d12.s1 Soares_placenta_8to9weeks_2NbHP8to9W Homo sapiens cDNA clone IMAGE:257399 3' similar to contains LTR7.b3 LTR7 repetitive element;
Top Hit Database Source	NT	IN	Ŋ	NT	L	NT	NT	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	L		Z	EST_HUMAN	NT	NT	TN	TN	EST_HUMAN	ΝT	NT	NT	LN L	N	LN T	FOR THE PARTY OF T	ESI LICIMAN	N	EST_HUMAN
Top Hit Acession No.	4502316 NT	4757915 NT	4757915 NT	AF083822.1	2.0E-53 M61873.1	4506962 NT	2.0E-53 AL163281.2	2.0E-53 AL163281.2		2.0E-53 BF334740.1				AB026898.1	1.0E-53 BF364201.1	9536.1	4507500 NT	4507500 NT	4508788 NT	BE386785.1	4504610 NT	4507848 NT	4507848 NT	4507848 NT	4507848 NT	6005700 NT	7 605030	7.0E-04-74612057.1		V27177.1
Most Similar (Top) Hit BLAST E Value	2.0E-53	2.0E-63	2.0E-53	2.0E-63 AI	2.0E-53				2.0E-53	2.0E-53			0.0	1.0E-53 A	1.0E-53			9.0E-54	9.0E-54	8.0E-54	8.0E-54	8.0E-54	8.0E-54	8.0E-54	8.0E-54	8.0E-54	100	7.00-04	7.0E-54	7.0E-54 N
Expression Signal	10.54	6.93	6.83	1.18	2.06	1.07	1.12	1.12	3.11	3.11	5.6	6.0			1.42	5.14	0.8	8.0	4.71	3.09	1.33	1.25	1.26	1.08	1.08	20.81	00	07:	1.54	4.61
ORF SEQ ID NO:		22446	22447	22933			24735	24736				21200								19988	21568	24307				25612		20100	21563	21945
Exan SEQ ID NO:	12365		12559	13131	13877	14288	14981	ш			17280	11334	70000	13283	15925	17129	15019	15019	19439	10172	11692	14517	14517	14517	14517	15529	7	2020	11687	12045
Probe SEQ ID NO:	2490	2694	2694	3207	3970	4380	5091	5091	5331	5331	7413	1429	7000	2000	6021	7252	. 6152	5152	5244	200	1794	4629	4620	5030	2030	5614	07.0	S/S	17889	2158

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wd26d11x1 Soares_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:2329269 3' similar to TR:002711 Homo sapiens similar to nuclear factor related io kappa B binding protein (H. sapiens) (LOC63182), mRNA qb67g03 x1 Soares_fetal_heart_NbHH19W Homo sapiens cDNA clone IMAGE:1705204 3' similar to 270112.r1 Soares_testis_NHT Homo sapiens cONA clone IMAGE:727727 5' similar to TR:G191315 Tupala belangeri beta-actin mRNA, partial cds i EST177696 Jurkat T-cells VI Homo sapiens cDNA 5' end similar to glyceraldehyde-3-phosphate Homo saplens BMX non-receptor tyrosine kinaise (BMX) mRNA al92c08.st Soares_parathyrcid_tumor_NbHPA Homo saplens cDNA clone IMAGE:1388270 3' al92c08.st Soares_parathyrcid_tumor_NbHPA Homo saplens cDNA clone IMAGE:1388270 3' DKFZp434E0731_r1 434 (synonym: htes3) Homo sapiens cDNA done DKFZp434E0731 5' Homo saplens killer cell lectin-like receptor subfamily G, member 1 (KLRG1), mRNA 602019408F1 NCI_CGAP_Brn67 Homo sapiens cDNA clone IMAGE:4155121 5' Homo sapiens hypothetical protein DKFZp434M036 (DKFZp434M035), mRNA Homo sapiens chloride channel 6 (CLCN9) mRNA Homo sapiens DNA for MICB, exon 4, 5 and partial cds Homo sapiens lymphocyte antigen 75 (LY75) niRNA, and translated products Homo sapiens lymphocyte antigen 75 (LY75) niRNA, and translated products EST185371 Colon carcinoma (HCC) cell line Homo sapiens cDNA 5' end G191315 ANDROGEN-DEPENDENT EXPRESSED PROTEIN.; **Top Hit Descriptor** ZINC FINGER PROTEIN 84 (ZINC FINGER PROTEIN HPF2) EST366629 MAGE resequences, MAGC Homo sapiens cDNA RC1-BT0313-131199-011-b09 BT0313 Homo sapiens cDNA RC3-ST0197-151099-011-f08 ST0197 Homo sepiens cDNA AV754746 TP Homo saplens cDNA clone TPGAAC10 5' Homo saplens nuclear antigen Sp100 (SP100) mRNA IL-BT189-190399-007 BT189 Homo sapiens cDNA 002711 PRO-POL-DUTPASE POLYPROTEIN Human mRNA for KIAA0077 gene, partial cds Human mRNA for KIAA0077 gene, partial cds H.sapiens shc pseudogene, p66 isoform H.saplens she pseudogene, p66 Isoform contains OFR.11 OFR repetitive elemen dehydrogenase EST HUMAN Top Hit Database Source HUMAN EST_HUMAN EST_HUMAN **EST HUMAN** EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN HUMAN EST_HUMAN EST_HUMAN SWISSPROT EST EST 눋 눋 Þ F 4505052 NT 8922148 NT z 4507164 NT 5031900 11417222 4505052 4502872 Top Hit Acession 3.0E-54 AW748965.1 2.0E-54 50319 2.0E-54 4507 AV754746.1 4.0E-54 AA306764.1 4.0E-54 D38521.1 3.0E-54 AA844061.1 3.0E-54 AA393362.1 3.0E-54 AW954559.1 6.0E-54 AB003618.1 6.0E-54 AW813567. 3.0E-54 AA844061.1 3.0E-54 BF345600.1 3.0E-54 AA313487.1 3.0E-54 AL110383.1 AI160189.1 4.0E-54 AF110103. ė 4.0E-54 A1935086.1 3.0E-54 AI908757.1 4.0E-54 D38521.1 8.0E-54 Y09846.1 6.0E-54 Y09846. P51523 8.0E-54 8.0E-54 8.0E-54 6.0E-54 6.0E-54 5.0E-54 7.06-52 Aost Similar (Top) Hit BLAST E Value 3.24 5.11 0.88 1.48 1.68 4.17 3.34 23 6.24 1.54 0.87 0.87 1.17 1.94 9.84 1.71 106.88 3.24 6.29 1.77 Expression Signal 27925 21606 22968 23612 28143 26394 26395 28565 28847 20374 19802 20712 21536 21537 21105 21605 19888 ORF SEQ 22287 Ö N O 18478 11730 14255 10865 1663 15512 16235 18563 18932 17681 10009 13169 13832 14658 17899 11993 11663 13093 10072 12396 12454 18235 19714 SEO ID 14658 10563 11248 10147 18309 ÿ SEQ ID 8675 9199 1342 8611 3248 4359 3168 9242 4774 8750 940 5598 6373 626 7831 1833 1833 2104 6373 3923 930 764 1764 8 2583 ÿ

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	_		_		,			_	_	,	_	_	, .	_	n	71	4		-		17:14	-#		ant.		40		-11 11
Top Hit Descriptor	nt78a09.s1 NCI_CGAP_Pr3 Homo sapiens cDNA clone IMAGE:1204600 similar to contains element L1 repetitive element;	au92g03.y/ Schneider fetal brain 00004 Homo sapiens cDNA clone IMAGE:2783764 5' similar to SW:CUL1_HUMAN Q13616 CULLIN HOMOLOG 1;	Homo sapiens chromosame 21 segment HS210010.	wy60b12x1 Soares_NSF_F8_9W_OT_PA_P_S1 Homo seplens cDNA done IMAGE:2552927 3' similar to TR:062084 Q62084 PHOSPHOLIPASE C NEIGHBORING;	nj45g09.s1 NCI_CGAP_Pr9 Homo sapiens cDNA clone IMAGE:995488 similar to gb:X53777 60S RIBOSOMAL PROTEIN L23 (HUMAN);	Homo sapiens chaperonin containing T-complex subunit 6 (CCT6) mRNA	Homo sapiens chromosome 21 segment HS21C001	Homo sapiens peptidylarginine delminase type III (LOC51702), mRNA	Homo sapiens dihydropyridine receptor alpha 2 subunit (CACNA2D1) gene, exon 7	Homo sapiens small inducible cytokine subfamily A (Cys-Cys), member 14 (SCYA14) mRNA	1243c11.y1 NOI_CGAP_Bm52 Homo sapiens cDNA clone IMAGE:2291348 5	Homo sapiens KIAA0100 gene product (KIAA0100), mRNA	Homo sapiens mRNA for KIAA1591 protein, partial cds	Homo sapiens mRNA for KIAA1591 protein, partial cds	Homo sapiens neurofibromin 1 (neurofibromatosis, von Recklinghausen disease, Watson disease) (NF1), mRNA	Homo sapiens mRNA for brain ryanodine receptor, complete cds	Homo saplens Janus kinase 2 (a protein tyrosine kinase) (JAK2), mRNA	Homo sapiens pescadillo (zebrafish) homolog 1, containing BRCT domain (PES1), mRNA	Homo sapiens period (Drosophila) homolog 3 (PER3), mRNA	601899230F1 NIH_MGC_19 Homo sapiens cCNA clone IMAGE:4128535 5'	AU077341 Sugano cDNA library Homo sapiens cDNA clone Zrv6C880 similar to 5'-end region of Human	gamma-glutamy transpeptidase mKNA, b end	Homo sapiens RFB30 gene for RING finger protein	Homo sapiens RFB30 gene for RING finger protein	fh02a02x1 NIH_MGC_17 Homo capiens cDNA clone IMAGE:2960907 6'	y/28e04.r1 Soares fetal liver spleen 1NFLS Horno saplens cDNA clone IMAGE:127998 5' similar to SP:C561 BOVIN P10897 CYTOCHROME:	ak28a11.s1 Soares_testis_NHT Homo saplens cDNA clone IMAGE:1407260 3'	AU139909 PLACE1 Homo sapiens cDNA clone PLACE1011576 5
Top Hit Database Source	EST_HUMAN	EST_HUMAN	LN	EST_HUMAN	EST HUMAN	LN	TN	FZ	NT	L	EST_HUMAN	LN	NT	LΝ	μN	N	LN L	FX	N	EST_HUMAN		ESI HUMAN	NT	NT	EST_HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN
Top Hit Acession No.	AA655008.1	AW163175.1	2.0E-54 AL163210.2	2.0E-54 AW057524.1	AA532925.1	4502642 NT	AL163201.2	7708448 NT	AF08382	4759069 NT	BE047864.1	11426657 NT	2.0E-64 AB046811.1	2.0E-54 AB046811.1	11426544 NT	4B001025.1	11429127 NT	7657454 NT	8567387 NT	3F315418.1		1.0E-54 AU077341.1	Y07829.2	8.0E-55 Y07829.2	8.0E-65 AW 409714.1	7.0E-55 R09348.1	4A889581.1	7.0E-55 AU139909.1
Most Similar (Top) Hit BLAST E Value	2.0E-54	2.0E-54	2.0E-54	2.0E-54	2.0E-54	2.0E-54	2.0E-54	2.0E-54	2.0E-54	2.0E-54	2.0E-54 B	2.0E-54	2.0E-64	2.0E-54	2.0E-54	2.0E-54 A	2.0E-54	2.0E-54	2.0E-54	1.0E-54 B		1.05-54	8.0E-55	8.0E-55	8.0E-55	7.0E-55	7.0E-55	7.0E-55
Expression Signal	9:	1.3	1.82	1.65	4.06	2.03	1.13	1.45	0.84	3.75	1.34	3.59	18.19	18.19	8.14	3.62	1.26	2.57	1.46	1.07		2.20	14.58	2.32	2.76	1.19	1.26	1.59
ORF SEQ ID NO:	21290	22261	22321	22587				24450	24790	25116	L	25503	26564	25565	26243	27617	17872		25216							92802		27393
Exan SEQ ID NO:	11434	12367	12428	12793	13418	14010	14240	14664	15023	15283	15360	15439	15488	15488	16093	17403	17599	18770	19244	14259	1,00	19340	11201	11204	18402	10981	17173	17191
Probe SEQ ID NO:	1529	2483	2556	2865	3501	4110	4343	4780	5156	5363	5440	5521	6223	5223	6227	7552	7749	8963	5693	4363	0.00	7005	1294	1297	8530	1085	7297	7315

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Probe SEQ (D NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
8541	18413	28679	12.75	7.0E-55]AI	A1561056.1	EST_HUMAN	tq29f09.x1 NCI_CGAP_Utf Homo sepiens cDNA clone IMAGE:2210249 3'
8541		28680	12.75		.1	EST_HUMAN	tq29f09.x1 NCI_CGAP_Ut1 Homo sapiens cDNA clone IMAGE:2210249 3'
9823			6.4.3			EST_HUMAN	ym57g07.r1 Soares infant brain 1NIB Homo sapiens cDNA clone IMAGE:52444 5'
8803		28908	1.98			ΙN	Homo sapiens mRNA for KIAA1501 protein, partial cds
1732	11633	21500	1.12		5.0E-55 AA704971.1	EST_HUMAN	zj95b09.s1 Soares_fetal_liver_spleen_1NFLS_IS1 Homo sapiens cDNA clone IMAGE:462617 3'
1732		21501	1.12		5.0E-55 AA704971.1	EST_HUMAN	zj95b09.s1 Soares_fetal_liver_splean_1NFLS_IS1 Homo sapiens cONA clone IMAGE:462617 3'
5941		25969	1.82		4502240 NT	NT	Homo sapiens arykulfatase E (chondrodysplasia punctata 1) (ARSE), mRNA
5941	15848	25970	1.82		4502240 NT	IN	Homo sapiens arysulfatase E (chondrodysplasia punctata 1) (ARSE), mRNA
7229		27296	2.06		4506302 NT	NT	Homo saplens protein tyrosine phosphatase, receptor type, alpha polypeptide (PTPRA) mRNA
0777	17620	27851	1.86			LΝ	Homo sapiens mRNA for KIAA0611 protein, partial cds
0777		27852	1.86		5.0E-55 AB014511.1	LΝ	Homo sapiens mRNA for KIAA0611 protein, partial cds
7869	17719	27965	1.19		5453765 NT	LN	Homo saplens nel (chicken)-like 2 (NELL2), mRNA
9283			2.13		11417972 NT	ΙN	Homo saplens pescadillo (zebrafish) homolog 1, containing BRCT domain (PES1), mRNA
49	12658	19843	1.36	4.0E-55	AW857994.1	EST_HUMAN	EST370064 MAGE resequences, MAGE Homo sapiens cDNA
656			33.95	4.0E-55	4826973 NT	TN	Homo sapiens RNA binding motif protein, Y chromosome, family 1, member A1 (RBMY1A1) mRNA
1422	11328		1.89	4.0E-55	7661713 NT	NT	Homo sapiens predicted osteoblast protein (GS3786), mRNA
1422	11328	21194	1.89		7661713 NT	NT	Homo sapiens predicted osteoblast protein (GS3786), mRNA
							7/52b10.x1 Soares_NSF_F8_9W_OT_PA_P_31 Homo saplens cDNA clone IMAGE:3390043 3' similar to
1498			1.26	4.0E-55	BF06141	EST_HUMAN	contains L1.t3 L1 repetitive element;
1979		21763	1.53	4.0E-55	4506180 NT	N	Homo saptens proteasome (prosome, macropain) subunit, alpha type, 2 (PSMA2) mRNA
1979		21764	1.53	4.0E-55		NT	Homo sapiens proteasome (prosome, macropain) subunit, alpha type, 2 (PSMA2) mRNA
2039		21824	7.73			L	Homo sapiens diacy/glycerol kinase, gamma (90kD) (DGKG) mRNA
2039		21825	7.73	4.0E-55	4503314 NT	NT	Homo sapiens diacylglycerol kinase, gamma (90kD) (DGKG) mRNA
2262		22048	1.25		4507784 NT	NT	Homo sapiens ubiquitin-conjugating enzyme E2 variant 1 (UBE2V1) mRNA
2546	12419		1.04			NT	Homo saptens Xq pseudoautosomal region; segment 1/2
3242		22984	1.38		AL163300.2	NT	Homo sapiens chromosome 21 segment HS21,C100
6857			6.44	4.0E-55	Ι₹I	LN	Homo sapiens chromosome 21 segment HS21/C010
8559	18429		4.46		W28189.1	EST_HUMAN	43c5 Human retina cDNA randomly primed sublibrary Homo sapiens cDNA
9200	18933		2.38		4.0E-55 BF303941.1	EST_HUMAN	601886575F2 NIH_MGC_17 Homo sapiens clinA clane IMAGE:4120338 5
. 9138			2.76		3.0E-55 BE178519.1	EST_HUMAN	PM1-HT0603-090300-001-g08 HT0603 Homo, saplens cDNA
9866	19356		1.65			NT	Homo saplens chromosome 21 segment HS21(C084
373		20160	2.3			Ŋ	Human endogenous retrovirus pHE.1 (ERV9)
539	10480					NT	Human endogencus retroviral DNA (4-1), complete retroviral segment
633	10570	20383	3.08	2.0E-55	4507296 NT	NT	Homo saplens syntaxin-binding protein 1 (STX3P1) mRNA, and translated products

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Table 4
Single Exon Probes Expressed in Heart

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Ѓор Hit Descriptor	Homo sapiens ubiquitin protein ligase E3A (human papilloma virus E6-associated protein, Angelman syndrome) (UBE3A) mRNA	CM1-HT0876-150800-357-g03 HT0876 Homo sapiens cDNA	am98h05.s1 Stratagene schizo brain S11 Horrio saplens cDNA clone IMAGE:1684185 3' similar to contains THR.b2 THR repetitive element;	AU119344 HEMBA1 Homo sapiens cDNA clone HEMBA1005583 5'	Homo sapiens mannose-6-phosphate receptor (cation dependent) (M6PR) mRNA	Oryctolagus cuniculus New Zealand white elorigation factor 1 alpha (Rabefla2) mRNA, complete cds	Homo sapiens mRNA for KIAA0903 protein, partial cds	601120116F1 NIH_MGC_20 Hamo sepiens cDNA clone IMAGE:2867027 57	601120116F1 NIH_MGC_20 Hamo sepiens cDNA clone IMAGE:2967027 5'	Homo sapiens SMA3 (SMA3), mRNA	Homo sapiens testis-specific Testis Transcript Y 1 (TTY1) mRNA, partial cds	Human mRNA for HLA-411E, a MHC class I molecule (major histocompatibility complex)	Homo sapiens mRNA for KIAA0408 protein, partial cds	Homo sapiens mRNA for KIAA0406 protein, partial cds	Homo sapiens CLP mRNA, partial cds	43c5 Human retina cDNA randomly primed sublibrary Homo sapiens cDNA	Homo saplens chromosome 21 segment HS21C067	Homo sapians chromosome 21 segment HS21C010	Homo sapiens DSCR5b mRNA, complete cdsi	Homo sapiens DSCR5b mRNA, complete cds	Homo sapiens hypothetical protein FLJ20126 (FLJ20126), mRNA	Homo sapiens hect domain and RLD 2 (HERC2), mRNA	Homo sapiens hect domain and RLD 2 (HERC2), mRNA	Homo sapiens chromosome 21 segment HS21C010	Homo saplens chromosome 21 segment HS21C010	Human infant brain unknown product mRNA, complete cds	Homo sapiens DNA-binding protein (LOC56242), mRNA	601237702F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3809552 5	yn62g03.r1 Soares adult brain N2b5HB55Y H¢mo sapiens cDNA clone IMAGE:173044 5' similar to contains THR repetitive element;	RC1-CT0252-231099-013-b07 CT0252 Homo saplens cDNA
Top Hit Database Source	L L	EST_HUMAN	EST_HUMAN	EST_HUMAN	MT	Ŋ	¥	EST_HUMAN	EST_HUMAN	NT	Ę	NT	TN	M	ᅜ	EST_HUMAN	NT	F	Ę	NT TN	NT	N	F	۲	E	Ę	ΤN	EST_HUMAN	EST_HUMAN	EST_HUMAN
Top Hit Acession	4507798	BE719986.1	A1002836.1		5060	U09823.1	AB020710.1	BE277861.1	BE277861.1	5803174 NT	AF000990.1	X13111.1	AB007866.2	AB007866.2	54057.1	W28189.1	AL163267.2	AL163210.2	AB037163.1	AB037163.1	8923125 NT	11433046 NT	11433046 NT	AL163210.2	AL163210.2	U50950.1	10567821 NT	BE379074.1	H19934.1	AW361213.1
Most Similar (Top) Hit BLAST E Value	2.0E-55	2.0E-55	2.0E-55	_		1.0E-55	1.0E-55		1.0E-55	1.0E-55	1.0E-55		1.0E-55	1.0E-55	1.0E-55	1.0E-55	1.0E-55	1.0E-55	1.0E-55	1.0E-55	1.0E-55	1.0E-55	1.0E-55	1.0E-55	1.0E-55	1.0E-55	1.0E-55	9.0E-56	7.0E-56	
Expression Signal	0.79	2.97	4.3	2.2	1.6	11.9	3.55	0.86	0.86	2.3	1.03	33.19	4.71	4.71	1.35	1.16	3.47	1.04	86.0	86'0	1.19	5.75	5.75	4.74	4.74	2.58	2.04	1.81	5.18	1.84
ORF SEQ ID NO:	22658			28446		19969	20888	21680	21681		22071	22238	22271	22272	22331	23082	23597		24392	24393	24727	25809	25810		28406		28959	26383	22457	26538
Exan SEQ ID NO:	12855		17160	18188	l	10154	11046	11802	11802	12161	12651	12346	12381	12381	12439	13282	13817	14097	14607	14607	14951	15700	15700	18163	18163	17894		16221	l	1
Probe SEQ ID NO:	2928	4665	7284	8319	91	182	1132	1907	1907	7722	2280	2470	2507	2507	2568	3363	3907	4197	4721	4721	5081	5794	5794	8284	8284	8745	8859	8368	2703	6504

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Table 4
Single Exon Probes Expressed in Heart

	Т	Т	т	1	_	_	_	τ-	Υ_	-	_	_	т	т-	Т	T	$\overline{}$	_	T -	_	Γ	ŕ			_	T	T-	`		Ë		ات
Top Hit Descriptor	RC1-C10252-231099-013-b07 CT0252 Homo saplens cDNA	RC3-BN0053-170200-011-h01 BN0053 Homo sapiens cDNA	43c5 Human retina cDNA randomly primed subilbrary Homo sapiens cDNA	CHR220038 Chromosome 22 exon Homo sepiens cDNA clone C22_55 5	Homo sapiens beta-tubulin mRNA, complete cds	Homo saplens beta-tubulin mRNA, complete cds	Homo sapiens tubulin, beta polypeptide (TUBB) mRNA	Homo saplens tubulin, beta polypeptide (TUBB) mRNA	Homo sapiens X-linked anhidroitic ectodermal dysplasia protein gene (EDA), exon 2 and flanking repeat	regions	Homo sapiens uncharacterized bone marrow protein BM031 mRNA, complete cds	Homo sapiens uncharacterized bone marrow protein BM031 mRNA, complete cds	Homo sapiens lymphocyte-specific protein 1 (LSP1) gene, LSP1-7 allele, partial cds	tm65g12.x1 NCI_CGAP_Bm25 Homo sapiens cDNA clone IMAGE:2163046 3'	tm65g12.x1 NCI_CGAP_Brn25 Homo sapiens cDNA clone IMAGE:2163046 3'	Homo sapiens hypothetical protein PRO1304 (PRO1304), mRNA	Homo sapiens oncogene TC21 (TC21), mRNA	EST28889 Cerebellum II Homo sapiens cDNA 5' end	EST28889 Cerebellum II Homo sapiens cDNA 5 end	Homo sapiens MHC class 1 region	Homo sapiens chromosome 21 segment HS21,0068	Homo sapiens superkiller viralicidic activity 2 (S. caravislae homolog)-like (SKIV2L), mRNA	Homo sapiens sparc/osteonectin, cwcv and kazal-tike domains proteoglycan (testican) (SPOCK) mRNA	Homo sapiens sparc/osteonectin, cwcv and kazial-like domains proteoolycen (testicen) (SPOCK) mRNA	Homo saplens lysosomal-associated membrans protein 2 (LAMP2), mRNA	Homo sapiens bone morphogenetic protein 5 (BMP5), mRNA	Homo sapiens KIAA0317 gene product (KIAA0317), mRNA	Homo sapiens mRNA, similar to rat myomegalin, complete cds	Homo sapiens nuclear pore complex Interacting protein (NPIP), mRNA	Homo sapiens nuclear pore complex interacting protein (NPIP), mRNA	Homo sapiens caveolin 3 (CAV3), mRNA	Homo sapiens caveolin 3 (CAV3), mRNA
Top Hit Database Source	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	۲	TN	Z	IN		L L	IN	LΝ	TN	EST_HUMAN	EST_HUMAN	IN	Z	EST_HUMAN	EST_HUMAN	ΙN	IN	INT	LN	LV	FN	Z,	NT	F	LN	NT	LN	NT
Top Hit Acesslon No.	AW361213.1	AW997712.1	W28189.1	H55099.1	AF141349.1	AF141349.1	4507728 NT	4507728 NT		AF003528.1	AF217508.1	AF217508.1	AF043349.1	A/498056.1	A1498086.1	8924029	6912697 NT	AA325826.1	AA325826.1	AF055066.1	AL163268.2	5902085 NT	4759163 NT	4759163 NT	11421124 NT	11418704 NT	11434956 NT	AB042556.1	5902013 NT	5902013 NT	11434876 NT	11434876 NT
Most Similar (Top) Hit BLAST E Value	7.0E-56	5.0E-56	5.0E-58	6.0E-56	4.0E-58	4.0E-56	4.0E-56	4.0E-58		4.0E-56	4.0E-56	4.0E-56	4.0E-56	4.0E-58	4.0E-58	3.0E-56	3.0E-58	3.0E-58	3.05-56	3.0E-56	3.0E-58	3.0E-56	3.0E-56	3.05-56	3.0E-56	3.0E-58	3.0E-56	3.0E-56	3.0E-56	3.0E-56	3.0E-56	3.0E-58
Expression Signal	1 8.	2.28	131	2.68	6.15	6.15	4.11	4.11		3.05	6.29	6.29	1.23	8.75	8.75	9.85	2.7	1.58	1.58	1.39	4.05	2.14	1.57	1.57	6.34	5.74	1.52	10.72	3.89	3.89	2.3	2.3
ORF SEQ ID NO:	26539	21434		24902	19805	19806	22431	22432		20268	25802	25803		28417	28418	21081	21892	22804	22805		24008	24154	25467	25468	28141	27185	28078	28259	28771	28772	26316	25316
Exam SEQ ID NO:	16363	11568	17790	19701	10012	10012	12541	12541		10457	15694	15694	17849	18174	18174	11225	11992	13013	13013	13674	14224	14364	15405	15405	16003	16994	17837	18012	18497		l	18955
Probe SEQ ID NO:	6504	1669	7940	9376	25	25	2676	2676		2781	5788	5788	7999	8296	8295	1318	2103	3086	3088	3761	4327	4470	5486	5488	6109	7117	7987	8124	8632	8632	9240	8240

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x05d10.x1 NCI_CGAP_Brn53 Homo sapiens cDNA clone IMAGE:2759251 3' similar to gb:U05875 INTERFERON-GAMMA RECEPTOR BETA CHAIN PRECURSOR (HUMAN); zq52a08.s1 Stratagene neuroepithelium (#937231) Homo sapiens cDNA clone IMAGE:645206 3 Homo saplens EphA4 (EPHA4) mRNA op67h02.s1 Soares_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:1581939 3 Human cGMP phosphodiesterase alpha suburit (CGPR-A) mRNA, complete cds Human cGMP phosphodiesterase alpha subunit (CGPR-A) mRNA, complete cds Macaca fascicularis protein tyrosine phosphatrise (PRL-1) mRNA, complete cds hg23c11.x1 NCI_CGAP_GC6 Homo sapiens cDN4 clone IMAGE:2946452 3* hg23c11.x1 NCI_CGAP_GC6 Homo sapiens cDN4 clone IMAGE:2946452 3* QV-BT077-130199-079 BT077 Homo sapiens cDNA zv51b12.r1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:757151 5 Homo sapiens SH3-domain binding protein 1 (SH3BP1), mRNA Homo sapiens smg GDS-ASSOCIATED PROTEIN (SMAP), mRNA Homo sapiens smg GDS-ASSOCIATED PROTEIN (SMAP), mRNA Homo sapiens hypothetical protein FLJ20371 (FLJ20371), mRNA QV4-ST0234-181199-037-f05 ST0234 Homo sapiens cDNA Homo sapiens hypothetical protein FLJ20371 (FLJ20371), mRNA Homo sapiens monocarboxylate transporter 3 (SLC16A8), mRNA Homo saplens SH3-domain binding protein 1 (SH3BP1), mRNA **Top Hit Descriptor** QV0-OT0033-070300-152-h03 OT0033 Homo sapiens cDNA Homo sapiens EphA4 (EPHA4) mRNA RC2-CT0163-220999-001-E02 CT0163 Homo sapiens cDNA RC4-BT0310-110300-015-f10 BT0310 Homo sepiens cDNA RC4-BT0310-110300-015-f10 BT0310 Homo sapiens cDNA AV703184 ADB Homo sapiens cDNA clone ADBCFG10 5' Homo sapiens aconitase 2, mitochondrial (ACO2), mRNA Homo sapiens mRNA for KIAA0960 protein, partial cds Homo saplens mRNA for KIAA1414 protein, partial cds Homo sapiens mRNA for KIAA0960 protein, pertial cds Homo sapiens mRNA for KIAA0837 protein, partial cds Homo saplens mRNA for KIAA0837 protein, partial cds Homo sapiens EphA4 (EPHA4) mRNA Homo sapiens EphA4 (EPHA4) mRNA EST HUMAN EST_HUMAN HUMAN **EST HUMAN** EST_HUMAN EST_HUMAN EST_HUMAN EST HUMAN HCMAN **EST HUMAN** HUMAN Top Hit Database Source EST EST F 눋 4758279 NT 8923349 NT 눋 눋 11545732 NT z 7657592 NT 7019528 4758279 8923340 11418185 Top Hit Acession 1.0E-56 AW589833.1 1.0E-56 AW589833.1 2.0E-56 BE064386.1 AW880885.1 4W818405.1 AW 264599.1 8.0E-57 AB023177.1 AA199818.1 2.0E-56 AV703184.1 AB020844.1 8.0E-67 AA971001.1 8.0E-57 AB020644.1 2.0E-56 AB037835.1 AA496109. 8.0E-67 AB023177. ė 1.0E-56 AF190930. 1.0E-58 AI905162.1 2.0E-56 M26061.1 2.0E-56 M26061.1 8.0E-57 8.0E-57 2.0E-56 9.0E-57 9.0E-57 8.0E-57 8.0E-57 8.0E-67 8.0E-57 8.0E-57 7.0E-57 9.0E-57 8.0E-57 8.0E-57 7.0E-57 (Top) Hit BLASTE Value 6.35 11.78 1.08 4.78 5.79 1.63 87.78 87.78 2.94 1.02 1.52 1.55 1.02 1-1 0.98 3.32 1.27 1.97 Expression Signal 22113 22684 25915 25251 22353 23210 23321 23322 24606 27863 23786 19790 20642 21651 23060 24750 25916 26594 25251 26595 19790 ORF SEQ 20477 20079 ÖΝÖ 13405 13536 19188 10455 12675 12215 12215 12886 14838 14009 14009 6666 10792 11673 13255 15794 16415 16415 19188 SEQ ID 14975 19624 15794 19097 12462 10887 10258 ġ Probe SEQ ID 2334 2959 716 2334 3489 9468 513 3622 4963 4109 3335 5207 5888 5888 8557 9607 9622 964 2 866 2592 609 4109 28 6107 6557 8771 ÿ

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Top Hit Descriptor	Homo saplens NME7 (NME7), mRNA	Homo sapiens NME7 (NME7), mRNA	Homo sapiens Kruppel-like factor 8 (KLF8), mRNA	Homo sapiens phosphatidylinositol 4-kinase 230 (pl4K230) mRNA, complete cds	Homo sapiens phosphatidylinositol 4-kinase 230 (pi4K230) mRNA, complete cds	Homo sapiens Xq pseudoautosomal region; segment 1/2	Homo saplens DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes, commiste ets)	601471226F1 NIH MGC 67 Homo sapiens cDNA clane IMAGE:3874135 5	Homo sapiens ubiquitin protein ligase E3A (human papilloma virus E6-associated protein, Angelman syndrome) (UBE3A) mRNA	nc13f07.s1 NCI_CGAP_Pr1 Homo saplens cDNA clone IMAGE:1008037 similar to SW:RS10_HUMAN P46783 40S RIBOSOMAL PROTEIN S10.;	EST64770 Hippocampus II Homo sapiens cDNA 5' end	7733b10.x1 NCI_CGAP_CLL1 Homo sapiens GDNA clone IMAGE:3298443 3' similar to WP:Y47H9C.2 CE20283 :	7733b10.x1 NCI_CGAP_CLL1 Homo sepiens cDNA clone IMAGE:3296443 3' similar to WP:Y47H9C.2	Home sarions celluline ts 4 201a chloride for current Inducer protein I/Cin) gene complete cds	RC3-CT0254-110300-027-d10 CT0254 Home saplens cDNA	601589896F1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:3944302 5'	42f6 Human retina cDNA randomly primed subilbrary Homo sapiens cDNA	Homo sapiens hypothetical protein FLJ11656 (FLJ11656), mRNA	Homo sapiens hypothetical protein FLJ11656 (FLJ11656), mRNA	AU117659 HEMBA1 Hamo sapiens cDNA done HEMBA1001910 5	2820473.5prime NIH_MGC_7 Homo sapiens cDNA clone IMAGE:2820473 5'	zb45d11.r1 Soares_fetal_lung_NbHL19W Horno sapiens cDNA clone IMAGE:306549 5'	RC0-HT0112-080999-001-C06 HT0112 Homo saplens cDNA	ak02b02.s1 Soares_parathyroid_tumor_NbHPA Homo sapiens cDNA clone IMAGE:1404747 3' similar to	contains Alu repstitive element contains element MER22 repetitive element ;	Homo sapiens chromosome 21 segment HS21C004	ye98h01.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:125809 5'	ye98h01.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:125809 6	Hamo sapiens chromosome 21 segment HS21,0083
Top Hit Database Source	TN	TN	TN	NT	TN	TN	L	EST HUMAN	늄	EST HUMAN	EST_HUMAN	EST HUMAN	NAM! H TAR		EST HUMAN	EST_HUMAN	EST_HUMAN	N7	TN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN		EST_HUMAN	NT	EST_HUMAN	EST_HUMAN	L
Top Hit Acession No.	7242158 NT	7242158 NT	E005979 NT	AF012872.1	AF012872.1	AJ271735.1	AROJAROR 1	BE783649.1	4507798 NT	AA230279.1	AA348335.1	BE676622.1	3E 678822 4	AE232708 1	_	BE796537.1	W28130.1	11545798 NT	11545798 NT	AU117659.1	4W248374.1	W23871.1	4W178575.1		AA845419.1	AL163204.2	207702.1	R07702.1	4L163283.2
Most Similar (Top) Hit BLAST E Value	7.0E-57	7.0E-57	7.0E-57	7.0E-57	7.0E-57	5.0E-57	4 0E-57	4.0E-57	3.0E-57	3.0E-57	3.0E-57			3.0E-57	3.0E-57	3.0E-57	3.0E-57	3.0E-57	3.0E-57	3.0E-57	3.0E-57	3.0E-57	3.0E-57		2.0E-67	2.0E-57	2.0E-57	2.0E-67	2.0E-57
Expression Signal	0.92	0.92	6.49	2.17	2.17	2.99	- 2	0.96	0.79	11.34	2.83		-	1 4	115.94	3.34	3.95	1.95	1.95	4.65	20.31	5.2	2.15		1.19	. 2.91	0.84	0.84	6.88
ORF SEQ ID NO:	22937		22956					24581			22119	22425		02220		25733		26817	26818	27212	28400	24907			22463		23223	23224	24086
Exan SEQ ID NO:	13136	13136	13157	13713	13713	19656	13608	1	10716	11215	12222	12535	10535	L		15630	16615	16830	16830	17019	18159	19721	19628	l				13421	14303
Probe SEQ ID NO:	3212	3212	3233	3801	3801	9934	3694	4935	787	1309	2342	2870	2870	3514	3639	5723	6736	6751	6751	7142	8279	9247	8096		2699	3392	3504	3504	4409

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Most Similar (Top) Hit Acession (Top) Hit Database Source Source	2.0E-57 AA016131.1 EST_HUMAN repetitive element;	1.29 2.0E-57 AF045452.1 NT Homo sapiens cell-line KG1 transcriptional regulatory protein p54 mRNA, complete cds	AF057722.1 NT	24084 NT	11424084 NT	1.12 1.0E-57 AW 503208.1 EST_HUMAN UI-HF-BN0-ekt-g-07-0-UI.r1 NIH_MGC_50 Homo sepiens cDNA clone IMAGE:3078348 5'	EST HUMAN		9.0E-58 BE395061.1 EST HUMAN	8.0E-58 BE868715.1 EST HUMAN		8.0E-58 AI798376.1	8.0E-58 11434921 NT		2.83 8.0E-58 7706132 NT Homo saplens DHHC1 protein (LOC51304), n/RNA	i L	7.0E-58 51/4542/NI	EST TOTAL	A DE RO DESOCROA 4 FOR LIMAN	6.0E-58 AU130689.1 EST HUMAN	1.19 6.0E-58 BE242150.1 EST HUMAN saplens cDNA clone TCAAP1219	1.19 6.0E-58 BE242150.1 EST HUMAN sapiens cDNA clone TCAAP1219	6.0E-58 11434746 NT	6.0E-58 11526291 NT
Expression (Top) Hit Signal BLAST E			L										L					ı	ı					
j W		 ۱	ত্র	lg:	8	21973			25238		20389	380	21588	589	T		.	67,007	2702	22105	22592	593	28000	1
ORF SEQ ID NO:		27084				L								L				1						
<u> </u>	5478 15398	7017 16894 2708	17515	18460	18460	12071	7045 16922	19055	19223	10512	10575	10575	11710	11710	12872	l	18112	6300 16179 2	10179	12205	2871 12798 2	2871 12798	17759	19109

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Top Hit Descriptor	Homo sapiens synaptojanin 1 (SYNJ1), mRNA.	RC4-NT0057-160600-016-b05 NT0057 Homo sapiens cDNA	CM3-UM0043-240300-127-e07 UM0043 Homo sapiens cDNA	CM3-UM0043-240300-127-e07 UM0043 Homo sapiens cDNA	CM3-UM0043-240300-127-e07 UM0043 Homo saplens cDNA	CM3-UM0043-240300-127-e07 UM0043 Homo sapiens cDNA	or88e07.s1 NCI_CGAP_Lu5 Homo sapiens dDNA clone IMAGE:1603908 3'	Homo sapiens placenta-specific 1 (PLAC1), mRNA	ym51h07.r1 Soares Infant brain 1NIB Homo sciplens cDNA clone IMAGE:52071 5'	Homo sapiens apical protein, Xenopus laevis-like (APXL), mRNA	Homo sapiens hypothetical protein FLJ10828 (FLJ10826), mRNA	Homo sapiens chromosome 21 segment HS21 C018	Homo sapiens cat eye syndrome chromosome region, candidate 1 (CECR1), mRNA	Homo sapiens Ran GTPase activating protein it (RANGAP1), mRNA	Homo sapiens ATP synthase, H+ transporting, mitochondrial F1 complex, O subunit (oligomycin sensitivity	conferring protein) (ATP50) mRNA	Homo sapiens interleukin 10 receptor, beta (IL:10RB), mRNA	Homo sapiens coagulation factor IX (plasma thromboplastic component, Christmas disease, hemophilia B) FEO) ∞BNA	Homo saniens i thiguilling-contingating RIR-domein enzyme APOLLON mRNA, complete cds	Himen beta-nime-selantin (1980) nene exm 3	Human mRNA, Xa terminal portion	Homo sapiens EGF-like repeats and discoidin I-like domains 3 (EDIL3), mRNA	Homo sapiens E1B-55kDa-assoclated protein 5 (E1B-AP5), mRNA	yg10e02.r1 Soares Infant brain 1NIB Homo sapiens cDNA clone IMAGE:31693 5'	Homo sapiens peptide YY (PYY) mRNA	602185789F1 NIH_MGC_46 Homo sapiens cDNA clone IMAGE:4309943 5'	602185789F1 NIH_MGC_45 Homo sapiens cDNA clone IMAGE:4309943 5'	AV712977 DCA Homo sepiens cDNA clone DCAAZG04 5'	Homo sapiens 5-aminolevulinate synthase 2 (ALAS2) gene, complete cds	ba08b07.y1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:2823733 5' similar to gb:X69391 60S	RIBOSOMAL PROTEIN L6 (HUMAN); gb:X81987 M.musculus mRNA for TAX responsive element binding protein (MOLISE).	Program (NOOC)	MICO_ 10 TOTAL WIND BEADING COINT CIOUR HWANCE, 550 1911 O
Top Hit Database Source		EST_HUMAN	EST_HUMAN	EST_HUMAN (Γ	EST_HUMAN		T_HUMAN															T_HUMAN		EST_HUMAN	HUMAN	EST_HUMAN	L'N		CCT LIMAN	Т	ESI_HUMAN
Top Hit Acession No.	4507334 NT	BE763984.1	AW 797948.1	AW797948.1	AW797948.1	AW 797948.1	AA988183.1	11496282	H23072.1	11421330 NT	8922693 NT	AL163218.2	11526293 NT	11418177 NT		4502302 NT	4504834 NT	7.02648	4303040			5031660	11424059 NT	R17879.1	4758981 NT	BF569848.1	BF569848.1	AV712977.1	AF068624.1		4 000000	DE-200332.1	BE907186.1
Most Similar (Top) Hit BLAST E Value	5.0E-58		5.0E-58	5.0E-58	5.0E-58	5.0E-58	5.0E-58	5.0E-58		5.0E-58	5.0E-58	5.0E-58	5.0E-58	5.0E-58		4.0E-58	4.0E-58	i c	_				4.0E-58		3.0E-58	3.0E-58	3.0E-58		2.0E-58			2.0E-30	2.0E-08
Expression Signal	2.9	5.87	4.47	4.47	2.89	2.89	3.56	2.21	5.86	1.45	6.77	1.58	3.26	2.49		17.97	1.58		1.24	0.0	44	2.11	7.54	1.17	2.34	2.91	291	1.39	8.16		90	1200	4.42
ORF SEQ ID NO:	20081		20932	20933			23002		25764		26721					20147			2777						21129			L	20698				24986
Exan SEQ ID NO:	10261	10626	11088	11088	11088	11088	13202	15378	15856	15800	16527	17879	18650	l		10325	L		1135/	20,00	13204	13594	18515	10291	11273	13068	13066	l	1			-1	19441
Probe SEQ ID NO:	297	693	1178	1178	1177	1177	3281	6468	5748	5894	6647	7829	9215	9878		369	779		1432		3283	3680	8651	332	1367	3141	3141	6002	925		-	1268	5288

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
5288	19441	25010	4.42	2.0E-58	2.0E-58 BE907186.1	EST_HUMAN	601499961F1 NIH_MGC_70 Homo saplens cDNA clane IMAGE:3901911 5
	1		-				am57e02.x1 Johnston frontal cortex Homo sapiens cDNA clone IMAGE:1539674 3' similar to WP:ZK328.1 CE05065 UBIQUITIN CONJUGATING ENZYME; RECOVERIN SUBFAMILY OF EF-HAND CALCIUM
5721	\perp			ŀ	2.0E-58 AI124874.1	EST_HUMAN	BINDING PROTEIN;
6246			2.76			Z	Homo sapiens endocytic receptor Endotos (TEVIOLOS) mKNA, complete das
6246	ı			١		LN	Homo sapiens endocytic receptor Endo180 (ENDO180) mKNA, complete cds
8123				-		EST_HUMAN	601890812F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:4131891 5
8332	18209	28459			1.1	EST_HUMAN	hm25f08.x1 NCI_CGAP_Thy4 Home sapiens cDNA clone IMAGE:3013671 3
202	10638	20463	0.93	1.0E-58		NT	Human complement component C5 mRNA, 3'end
1052	10969	20811	5.45	1.0E-58	6274549 NT	TN	Homo sapiens NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 9 (22kD, B22) (NDUFB9), mRNA
1305	11212	21067	2.17	1.0E-58	1.0E-58 AW957182.1	EST_HUMAN	EST369252 MAGE resequences, MAGD Homb saplens cDNA
1305			2.17	1.0E-58	_	EST_HUMAN	EST369252 WAGE resequences, MAGD Homb sapiens cDNA
1375	11281	21136	1.07	1.0E-58	1.0E-58 AJ238093.1	NT	Homo sapiens partial AF-4 gene, exons 2 to 7 and Alu repeat elements
1641	11545		2.02	1.0E-58	BE465132.1	EST HUMAN	hy10f08.x1 NCI_CGAP_GC8 Homo sapiens cIDNA clone IMAGE:3196935 3'
2771	12633			1.0E-58	4759169 NT	NT	Homo sapiens sterol regulatory element binding transcription factor 2 (SREBF2) mRNA
3493		23215			4758081 NT	L	Homo sapiens chondroitin sulfate proteoglycan/2 (versican) (CSPG2) mRNA
3493					4758081 NT	NT	Homo sapiens chondroitin sulfate proteoglycan 2 (versican) (CSPG2) mRNA
3659	13573		48.0	1.0E-58	4507628 NT	LΝ	Homo sapiens transition protein 1 (during historie to protamine replacement) (TNP1) mRNA
4627	14515			1.0E-58	95963	NT	Human prohormone converting enzyme (NEC2) gene, exon 4
4895		24553	4.86	1.0E-58 A	A1141063.1 EST_!	EST_HUMAN	oz43h01.x1 Soares_NHIMPu_S1 Homo saplens cDNA clone IMAGE:1678129 3'
7146	17023			1.0E-58		ΝΤ	Homo sapiens myomesin (M-protein) 2 (165kD) (MYOM2), mRNA
8982			3.46		X63392	LN L	H.sapiens immunoglobulin kappa light chain variable region L14
2182	12069		27.47		4507378	TN	Homo saplens TATA box binding protein (TBP) mRNA
6229	16638	26826			761963.1	EST_HUMAN	wh50d06.x1 NCI_CGAP_Kid11 Homo sapiens CDNA clone IMAGE:2384171 3'
171	12660		1.63			EST_HUMAN	601458531F1 NIH_MGC_66 Homo sapiens cDNA clone IMAGE:3862086 6'
3088	13015	22807	6.21	5.0E-59		EST_HUMAN	wf48c11.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2358839 3'
4559	14451	24237	5.85	5.0E-59	5.0E-59 X83497.1	NT	H.saplens DNA for ZNF80-linked ERV9 long terminal repeat
							au66c07.x1 Schneider fetal brain 00004 Homolsapiens cDNA clone IMAGE:2781228 3' similar to contains
6173	15130	24850		_	1	EST_HUMAN	element TAR1 repetitive element;
7596	17447	27662	1.71	5.0E-59	5.0E-59 AV762869.1	EST_HUMAN	AV762869 MDS Homo saplens cDNA clane MDSEIC12 5'
8278			2.8		11434908 NT	TN	Homo sapiens hypothetical protein (LOC57143), mRNA
776	1		2.42		D80006.1	NT	Human mRNA for KIAA0184 gene, partial cds
4685	14571	24368	1.2	4.0E-59	4506758 NT	Į.	Homo sapiens ryanodine receptor 3 (RYR3) mRNA

PCT/US01/00666

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	7.7		Т	Т		Г			Г	П	Г	П		Г	П	П			Г		ΓΞ	Ė		<u> </u>	Т	Ť	7		ΤĒ	Γ	T	T
Top Hit Descriptor	Homo sapiens ryanodine receptor 3 (RYR3) mRNA	Homo sapians 17-bata-hydroxysteroid dehydrogenase IV (HSD17B4) gene, promoter region and exon 1	EST377582 MAGE resequences, MAGI Homo sapiens cDNA	Homo sapiens KIAA0680 gene product (KIAA0680), mRNA	Homo sapiens plasminogen activator, tissue (PLATa) mRNA	Homo sapians plasminogen activator, tissue (PLATa) mRNA	Homo sapiens mRNA for KIAA112 protein, partial cds	Homo sapiens mRNA for KIAA1112 protein, partial cds	Homo sapiens A kinase (PRKA) anchor protein 1 (AKAP1), mRNA	Homo sapiens A kinase (PRKA) anchor protein 1 (AKAP1), mRNA	Homo sapiens zona pellucida glycoprotein 2 (sperm receptor) (ZP2) mRNA	Homo sapiens Testis-specific XK-related protein on Y (XKRY) mRNA	Homo sapiens protein tyrosine phosphatase, receptor type, T (PTPRT), mRNA	Homo sapiens hypothetical protein PRO1741 (PRO1741), mRNA	Homo sapiens nuclear receptor co-repressor 1 (NCOR1), mRNA	Human mRNA for dbl proto-oncogene	Human mRNA for dbl proto-oncogene	Homo sapiens gamma-glutamyftransferase-like activity (GGTLA1), mRNA	Homo sapiens gamma-glutamytransferase-like activity 1 (GGTLA1), mRNA	EST180633 Jurkat T-cells V Homo sapiens cDNA 5' end	RC0-NT0036-100700-032-a07 NT0036 Homo sapiens cDNA	fh07h04x1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:2961654 5'	fh07h04.x1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:2961654 5'	wa38c12x1 NCI_CGAP_Kid11 Homo sapiens/cDNA clone IMAGE:2300182 3' similar to TR:Q86542 O86542 RTVI .H PROTEIN "contains I TR7 ki i TR7 renetitive element"	Approximate substitution me MAIA commission in the commission of t	nomo sapiens aipne-mount mivvy, compiere cas	601176757F1 NIH_MIGC_17 Homo sapiens cone IMAGE:3531927 5	oa56h11.s1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:1309029 3' similar to TR:Q13537 Q13537 MER37 TRANSPOSABLE ELEMENT, COMPLETE CONSENSUS SEQUENCE.	Homo sapiens mRNA for transcription factor	Homo sapiens zinc finger protein 275 (ZNF275), mRNA	Homo sapiens mRNA for transcription factor	Homo sapiens small nuclear ribonucleoprotein D3 polypeptide (18kD) (SNRPD3) mRNA
Top Hit Database Source	Ŀ	M	EST HUMAN	NT L	TN	IN	NT	N	٦	N	NT	Ę	Ę	۲	Į.	۲	NT	IN	N L	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	FOT HIMAN	NT CINCLE	- 2	EST_HUMAN	EST HUMAN	NT	F	NT	TN
Top Hit Acession No.	4506758 NT	AF057720.1	AW965524.1	7682247 NT	4505860 NT	4505860 NT	AB029035.1	AB029035.1	4502014 NT	4502014 NT	4508044 NT	4759329 NT	7427522 NT	8924074 NT	5454137 NT	X12558.1	X12556.1	11417868 NT	11417868 NT	AA309774.1	BF365554.1	AW410698.1	AW410698.1	A1831800 4	١		BE296411.1	AA748468.1		11419630 NT	AJ130894.1	4759159 NT
Most Similar (Top) Hit BLAST E Value	4.0E-59	4.0E-59	3.0E-59	3.0E-59	3.0E-59	3.0E-59	3.0E-59	3.0E-59	3.0E-59	3.0E-59	3.0E-59	3.0E-59	3.0E-59	3.0E-59	3.0E-59	3.0E-59	3.0E-59	3.0E-59	3.0E-59	2.0E-69	2.0E-59	2.0E-59 /	2.0E-59	2 05-50			1.0E-59	1.0E-59/		1.0E-59	1.0E-59 /	8.0E-60
Expression Signal	12	2.18	6.96	4.12	9.87	9.87	7.68	7.68	3.71	3.71	1.17	1.07	1.85	2.03	1.82	1.23	1.23	1.64	3.9	5.01	2.47	1.84	1.84	44.4	27.5	2.73	3.58	2.46	1.29	1.22	8.32	2.71
ORF SEQ ID NO:	24369			20002	21455	21466	21866		22811	22812	23449	24368	24406	25786	26377	26697	26698					28341	28342	25350	1				26488	27524		21224
Exan SEQ ID NO:	14571	19586	9668	10191	11584	11584	11972	11972	13017	13017	13666	14569	14619	15679	16215	16509	16509	19014	19101	17408	17853	18089	18089	18053	40844	1981	10131	12446	16322	17318	16322	11360
Probe SEQ ID NO:	4685	9356	0	221	1682	1682	2082	2082	3090	3090	3753	4683	4734	5772	6352	6829	6629	9333	9474	7555	8003	8205	8208	9038	2750	8	157	2575	6463	7400	8229	1455

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		RTP) mRNA	RTP) mRNA													205087 5' similar to contains		205087 5' similar to contains		201953 5' similar to contains	2123	12123'	:3078348 5'	3078348 5'	ated pol						milar to SW:UDP_MOUSE		
olligie Eauli rioues Expressed ili neali	Top Hit Descriptor	Homo saplens differentiation-related gene 1 (nickel-specific induction protein) (RTP) mRNA	Homo sapiens differentiation-related gene 1 (nickel-specific induction protein) (RTP) mRNA	Homo sapiens mRNA for KIAA1081 protein, partial cds	Human mRNA for integrin alpha-2 subunit	Homo sapiens S-antigen; retina and pineal gland (arrestin) (SAG), mRNA	Homo saplens KIAA0433 protein (KIAA0433), mRNA	Homo saplens KIAA0433 protein (KIAA0433), mRNA	Homo saplens chromosome 21 segment HS21 C004	Homo sapiens chromosome 21 segment HS21 C004	Homo saplens MHC class 1 region	Homo sapiens MHC class 1 region	Homo sapiens interleukin 10 receptor, beta (IL 10RB), mRNA	Homo sapiens cullin 4A (CUL4A) mRNA, complete cds	Homo sapiens ornithine decarboxylase 1 (ODC1) mRNA	yr12f04.11 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:205087 5' similar to contains	LTR5 repetitive element ;	yr12f04.r1 Soares fetal liver spleen 1NFLS Hano saplens cDNA clone IMAGE:205087 6' similar to contains	LTR5 repetitive element ;	yq78h09.r1 Soares fetal liver spleen 1NFLS Homo saplens cDNA clone IMAGE:201953 5' similar to contains OFR repetitive element:	wf52c07.x1 Soares NFL T GBC S1 Homo sapiens cDNA clone IMAGE:23592123	wf52c07.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2359212 3	UI-HF-BN0-akt-g-07-0-UI.r1 NIH_MGC_60 Homo sapiens cDNA clone IMAGE:3078348 61	UI-HF-BN0-akt-g-07-0-UI.r1 NIH_MGC_50 Hcmo sepiens cDNA clone IMAGE:3078348 5	EST11498 Uterus Homo sapiens cDNA 5' end similar to similar to retrovirus-related pol	601336446F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3690395 5'	601336446F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3690395 5'	Homo saplens prohibitin (PHB) mRNA	Homo sapiens Xq pseudoautosomal region; segment 1/2	RC3-LT0023-200100-012-a01 LT0023 Homo sapiens cDNA	oleohtt. ys NCI_CGAP_Kid3 Homo sapiens cloNA clone IMAGE:1534053 5' similar to SW:UDP_MOUSE P52824 URIDINE PHOSPHORYLASE	Homo saplens proline dehydrogenase (proline (xidase) (PRODH) mRNA	
אום בייחוו הוחחם	Top Hit Database Source			NT TN	TN TN				H.	NT IN	님	T.		Ŧ.		T	EST_HUMAN L		EST_HUMAN L	FST HIMAN C	Т	HUMAN	П			EST_HUMAN 6	Γ		IN IN	EST_HUMAN R	O PET HIMAN	t	
Sillo Sillo	Top Hit Acession No.	5174656 NT	5174656 NT	8.0E-60 AB029004.1	X17033.1	11428949 NT	11417118 NT	11417118 NT	8.0E-60 AL163204.2				4504634 NT	AF077188.1	4505488 NT		H58041.1		H58041.1	H52456 1	1807917.1		4.0E-60 AW 503208.1	4.0E-60 AW 503208.1	4.0E-60 AA299037.1	3.0E-60 BE562611.1		11190	3.0E-60 AJ271735.1	3.0E-80 AW838188.1	3 0F-60 A 1782814 1	5174644 NT	
	Most Similar (Top) Hit BLAST E Value	8.0E-60	8.0E-60	8.0E-60	8.0E-60	8.0E-60	8.0E-60	8.0E-60	8.0E-60	8.0E-60	7.0E-60	7.0E-60	7.0E-60	7.0E-60	7.0E-60		7.0E-60 H		7.0E-60 H	6 0F-60 H	5.0E-60	6.0E-60	4.0E-60	4.0E-60	4.0E-60	3.0E-60	3.0E-80	3.0E-60	3.0E-60	3.0E-80	3.05-60	3.0E-60	
	Expression Signal	2.7	2.7	1.41	2.6	2.26	1.68	1.68	5.38	5.38	69'9	32.94	1.15	1,56	2.63		3.28		1.96	7.13	192	1.94	0.93	0.93	1.12	4.85	4.85	2.22	2.08	2.08	131	5.4	
	ORF SEQ ID NO:	21911	21912	L		27240	27465	27466	28344	28345	20504	20504	20567	21865	23765		27488		28843		19878	19879							24035	25443	92876		
	Exan SEQ ID NO:	12013	12013	15553	16524	17051	17260	17260	18091	18091	10669	10669	10727	11971	13988		17279		18559	16792	1	10062	12075	12075		11712	11712	11723	14250	15383	15111		
	Probe SEQ ID NO:	2125	2125	5640	6644	7174	7451	7451	8207	8207	737	738	798	2081	4088		7412		8671	6914	28	78	2188	2188	2942	1816	1815	1826	4324	5463	6153	9689	

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Top Hit Descriptor	ox66d09.x1 Soares_NhHMPu_S1 Homo sepiens cDNA clone IMAGE:1660337 3' similar to SW:FORM_MOUSE Q05860 FORMIN;	Homo sapiens proline dehydrogenase (proline oxidase) (PRODH) mRNA	N 601646227F1 NIH MGC 60 Homo sapiens cDNA clone IMAGE:3930990 5'	Homo sapiens solute carrier (SLC25A18) mRNA, complete cds; nuclear gene for mitochondrial product	H.sapiens 41kDa protein kinase related to rat ERK2	Human bor protein mRNA, 5' end	Homo sapiens chromosome 21 unknown mRNA	Home sapiens pro-alpha 2(I) collagen (COL1A2) gene, complete cds	Homo sapiens corticotropin releasing hormone receptor 2 (CRHR2) mRNA	Homo sapiens corticotropin releasing hormone receptor 2 (CRHR2) mRNA	N EST181949 Jurkat T-cells V Homo sapiens cDNA 5' end similar to similar to prothymosin, alpha	IN EST181949 Jurkat T-cells V Homo sapiens cD/NA 5' end similar to similar to prothymosin, alpha	Human pre-B cell stimulating factor homologue (SDF1b) mRNA, complete cds	Homo sapiens sema domain, transmembrane domain (TM), and cytoplasmic domain, (semaphorin) 6A (SEMA6A), mRNA	Homo saplens sema domain, transmembrane domain (TM), and cytoplasmic domain, (semaphorin) 6A (SEMA6A), mRNA	Homo saplens non-histone chromosome protein 2 (S. cerevislae)-like 1 (NHP2L1), mRNA	Homo saplens somatostatin receptor subtype 3/(SSTR3) gene, 5' flanking region and partial cds	Homo saplens similar to HSPC022 protein (H. saplens) (LOC63504), mRNA	Homo sapiens gene for AF-6, complete cds	Homo septens calcium channel, voltage-dependent, alpha 11 subunit (CACNA11), mRNA	NN PM3-HT0605-270200-001-e06 HT0605 Homo sapiens cDNA	N AU143389 Y79A41 Homo sapiens cDNA clone Y79AA1001854 6	Homo sapiens chromosome 21 segment HS21,0086	Г			IN AU119344 HEMBA1 Homo sapiens cDNA clone HEMBA1005583 5'			Human endogenous retrovirus pHE.1 (ERV9)
Top Hit Database Source	EST_HUMAN	N	EST_HUMAN	۲	Z F	N	NT	N.	ΝΤ	LN	EST_HUMAN	EST_HUMAN	NT	NT	ΤŃ	N L	NT	LN	N.	ΝΤ	EST_HUMAN	EST_HUMAN	LΝ		EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	ΝΤ
Top Hit Acession No.	A1040235.1	5174644 NT	BF102612.1	AY008285.1	211694.1	M24603.1	AF231919.1		4503044 NT	4503044 NT	AA311159.1	AA311159.1	L36033.1	11991659 NT	11991659 NT	11418192 NT	AF068757.1	11418068 NT	AB011399.1	11418157 NT	BE178586.1	AU143389.1	AL163285.2		~	•	AU119344.1		AW006478.1	X57147.1
Most Similar (Top) Hit BLAST E Value	3.0E-60	3.0E-60	3.0E-60	2.0E-80	2.0E-60	2.0E-60	2.0E-60	2.0E-60	2.0E-60	2.0E-60	2.0E-60	2.0E-60	2.0E-80	2.0E-60	2.0E-80	2.0E-60	2.0E-60	2.0E-60	2.0E-60	2.0E-60	1.0E-60	1.0E-60	1.0E-60		1.0E-80	1.0E-60	9.0E-61	8.0E-61	8.0E-61	8.0E-61
Expression Signal	2.59	4.7	3.84	1.79	2.89	- 83.	0.78	1.57	2.44	2.44	3.22	3.22	3.86	1.89	1.89	2.86	1.31	1.46	1.47	1.4	1.56	1.12	1.1		2.9	1.58	1.9	1.39	1.39	1.74
ORF SEQ ID NO:	27061	27147	27637	19810	21171	21482	23543			24881	26238	26239	27216	27808	27807						20264		24533						22396	
Exan SEQ ID NO:	16867	16954	17420	10015	11310	11593	13750	15816	15103	15103	16088	16088	17022	17582	17582	19123	19573	19209	19220	19420	10453	13739	14754	l		16978	10999	H	12502	12848
Probe SEQ ID NO:	0669	7077	7669	88	1405	1691	3839	5910	6093	8093	6222	6222	7145	7732	7732	9509	6296	9641	8998	8957	511	3827	4874		7086	7101	1083	2635	. 2635	2921

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	-						
Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Tôp Hit Descriptor
122	10098	19918	0.94	7.0E-61	7706670 NT	LN	Homo sapiens PXR2b protein (PXR2b), mRNA
122	10098	18919	0.94	7.0E-61	IN 0289077	NT	Homo saplens PXR2b protein (PXR2b), mRNA
123	10098	19918	0.86	7.0E-61	TN 0788077	TN	Homo saplens PXR2b protein (PXR2b), mRNA
123	10098	19919	0.86	7.0E-61	TN 0789077	LN	Homo sapiens PXR2b protein (PXR2b), mRNA
265			2	6.0E-61	BE409310.1	EST_HUMAN	601300938F1 NIH_MGC_21 Homo saplens cDNA clone IMAGE:3635480 5'
797		20564	1.69	6.0E-61	BE409310.1	r_HUMAN	601300938F1 NIH_MGC_21 Homo sepiens cDNA clone IMAGE:3635480 5
1289	11208	21060	10.28	6.0E-61	AF119860.1	NT	Homo sapiens PRO2014 mRNA, complete cds
1812		21376	0.95	6.0E-61			601109238F1 NIH_MGC_16 Homo sapiens cDNA done IMAGE:3350145 5'
1628	11532	21392	2.12		AA596033.1	EST_HUMAN	nn66h09:s1 NCI_CGAP_Lar1 Homo sapiens dJNA clone IMAGE:1088897 3'
3266	13189	22887	8.19	6.0E-61	AU130689.1	EST_HUMAN	AU130889 NT2RP3 Homo sapiens cDNA clonis NT2RP3001263 5'
5674	15583	25684	2:92	6.0E-81	S79249.1	NT	ig-beta/B29=CD79b (alternatively spliced) [human, B cells, mRNA Partial, 375 nt]
6344	16207	26370	1.93	6.0E-61		LN	Human autosomal dominant polycystic kidney disease protein 1 (PKD1) gene
6492		126521	2.03	6.0E-61	AF035737.1	NT	Homo sapiens general transcription factor 2-I (GTF2I) mRNA, complete cds
9417	10723	20564	1.43	6.0E-81	BE409310.1	EST_HUMAN	601300938F1 NIH_MGC_21 Hamo sapiens cDNA clone IMAGE:3635480 5
1655	11558	21421	1.78	5.0E-61	4506008 NT	NT	Homo saplens protein phosphatase 1, regulatory subunit 10 (PPP1R10) mRNA
3000	12928	22720	1.92	6.0E-61	AL163279.2	TN	Homo sapiens chromosome 21 segment HS21C079
3114	13039	22835	0.84	5.0E-61	AB020832.1	NT	Homo sapiens mRNA for KIAA0825 protein, partial cds
3161	13086	22890	1.9	5.0E-61	4502166 NT	Ę	Homo sapiens amyloid beta (A4) precursor protein (protease nexin-il, Alzheimer disease) (APP), mRNA
3899			1.68	5.0E-61	AJ229041.1		Homo sapiens 959 kb cantig between AML1 end CBR1 on chromosome 21q22; segment 1/3
9213	18941		2.76	4.0E-61	AV731140.1		AV731140 HTF Homo sapiens cDNA clone HTFARB01 5'
4118	14019	23797	1.13	3.0E-61	BE396279.1	EST_HUMAN	601309785F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3631220 5'
490			1.5	2.0E-61	8922829	۲	Homo sapiens hypothetical protein FLJ11026 (FLJ11026), mRNA
1194	11104		1.35	2.0E-61	BE168410.1	EST_HUMAN	QV3-HT0513-060400-147-d01 HT0513 Homo sapiens cDNA
1184	11104	20951	1.35	2.0E-61	BE168410.1	EST_HUMAN	QV3-HT0513-060400-147-d01 HT0513 Homo sapiens cDNA
				L			ys33d11.s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:246453 3' similar to
1042	\perp		1.31	2.0E-01	T.ASOSCA	ES HOWAN	gb:L29444 005 NIBOSOWIAL FNO I EIN L29A((TOWAN))
2108		76917	-		c000c/4		ווטוווס מקומום מחווס וו (סבסוג) ווויינים
2604	12472		1.18	2.0E-61	N39397.1	EST_HUMAN	yy03111.r1 Soares metanocyte 2NbHM Homo sapiens cDNA clone IMAGE:270189 5
5871	15777	25896	1.7	2.0E-61	11426166 NT		Homo sapiens ATPase, H+ transporting, lysosomal (vacuolar proton pump) non-catalytic accessory protein 1A (110/116kD) (ATP6N1A), mRNA
7212	17089	27279	1.33	2.0E-61	AV694317.1	EST_HUMAN	AV694317 GKC Homo saplens cDNA clone GICCELG06 5'
7707	17557		1.62	2.0E-81	AW 500256.1	T HUMAN	UI-HF-BN0-ekd-f-12-0-Ut.rt NIH_MGC_50 Homo saptens oDNA clone IMAGE:3076774 5'
7885	17735	27979	3.09	2.0E-61	11421778	LN.	Homo saplens polymerase (RNA) III (DNA directed) (39kD) (RPC39), mRNA

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Table 4
Single Exon Probes Expressed in Heart

Top Hit Descriptor	NUCLEOLAR TRANSCRIPTION FACTOR 1 (UPSTREAM BINDING FACTOR 1) (UBF-1) (AUTOANTIGEN NOR-90)	qg56a04.x1 Soares_testis_NHT Homo seplens cDNA clone IMAGE:1839150 3' similar to TR:O15103 O15103 HYPOTHETICAL 27.3 KD PROTEIN ;	Human zinc finger protein ZNF131 mRNA, partial cds	Homo sapiens CGI-56 protein (CGI-56), mRNA	wi04d02.x1 NCI_CGAP_CLL1 Homo saplens cDNA clone IMAGE:2389251 3'	wi04d02.x1 NCI_CGAP_CLL1 Homo sapiens cDNA clone IMAGE:2389261 3'	Homo sapiens CGI-18 protein (LOC51008), mRNA	MR3-ST0203-130100-025-a09 ST0203 Homo sapiens cDNA	wx61e07.x1 NCI_CGAP_Lu28 Home sapients cDNA clone IMAGE:2547204 3' similar to SW:GG95_HUMAN Q08379 GOLGIN-95, ;contains element MER22 repetitive element;	Homo saplens Xq pseudoautosomal region; segment 1/2	Homo sapiens Xq pseudoautosomal region; segment 1/2	Homo sapiens ryanodine receptor 3 (RYR3) mRNA	zw78609.s1 Soares_tests_NHT Homo saplens cDNA clone IMAGE:782344 3' similar to SW:NRDC_RAT P47245 NARDILYSIN;	fh07g09.x1 NIH_MGC_17 Homo sapiens cDNA clane IMAGE:2861616 51	Homo sapiens muscle specific gene (M9), mfRNA	Homo sapiens muscle specific gene (M9), mRNA	au71403.y1 Schneider fetal brain 00004 Homb sapiens cDNA clone IMAGE:2781701 6' similar to gb:M37104 ATP SYNTHASE COUPLING FACTOR 6, MITOCHONDRIAL PRECURSOR (HUMAN);	au71d03.y1 Schnelder fetal brain 00004 Homb sapiens cDNA clone IMAGE:2781701 5' similar to gb:M37104 ATP SYNTHASE COUPLING FACTOR 6, MITOCHONDRIAL PRECURSOR (HUMAN);	au71d03.y1 Schneider fetal brain 00004 Homo saplens cDNA clone IMAGE:2781701 5' similar to gb:M37104 ATP SYNTHASE COUPLING FACTOR 6, MITOCHONDRIAL PRECURSOR (HUMAN);	au71d03.y1 Schneider fetal brain 00004 Homo sapiens cDNA clone IMAGE:2781701 5' similar to gb:M37104 ATP SYNTHASE COUPLING FACTOR 6, MITOCHONDRIAL PRECURSOR (HUMAN);	EST182043 Jurkat T-cells V Homo sapiens cDNA 5' end	(wf12b08.x1 Soares_NFL_T_GBC_S1 Homo sepiens cDNA clone IMAGE:2350359 3' similar to gb:X57138_ma1 HISTONE H28.2 (HUMAN);	wf12b08.x1 Sogres_NFL_T_GBC_S1 Homo seplens cDNA clone IMAGE:2350359 3' similar to gb:X57138_ma1 HISTONE H2B.2 (HUMAN);	Homo sapiens keratin 18 (KRT18) mRNA
Top Hit Database Sœuroe	SWISSPROT	EST_HUMAN	N	NT	EST_HUMAN	EST_HUMAN	IN	EST_HUMAN	EST HUMAN	N-	N	N	EST HUMAN	EST HUMAN	님	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	NT
Top Hit Acession No.	P17480	AI208681.1	U09410.1	11418255 NT	AI762801.1	AI762801.1	11431139 NT	AW814393.1	A1950528.1	AJ271735.1	AJ271735.1	4506758 NT	AA431093.1	AW410687.1	11425574 NT	11425574 NT	AW161479.1	AW 161479.1	AW 161479.1	AW 161479.1	AA311281.1	AI827900.1	A1827900.1	4557887 NT
Most Similar (Top) Hit BLAST E Value	7.0E-62	7.0E-62	8.0E-62	6.0E-82	6.0E-62	6.0E-62	6.0E-62	6.0E-62	5.0E-62	5.0E-62			5.0E-62)	5.0E-62	5.0E-82	5.0E-62	4.0E-62	4.0E-62	4.0E-62	4.0E-62	4.0E-62	4.0E-62	4.0E-62	4.0E-62
Expression Signal	0.78	4	1.07	3.93	3.33	3.33	1.4	2.78	1.49	3.26	3.26	2.17	1.65	6.17	4.91	4.91	3.47	3.47	4.63	8.4	0.89	1.39	1.39	6.8
ORF SEQ ID NO:	23184	28829				26526	26873		20183		22133		23907				20597	20598	20597	20698		22183	22184	
Exan SEQ ID NO:	13378	18546	12896		16355			17265	10356	1	12236	ı	14131		18455	18455	10750	10750	10750	10750	11351	12286	12286	
Probe SEQ ID NO:	3462	8657	2969	3338	6496	6496	6804	7386	410	2356	2356	3372	4233	7482	8587	8587	822	822	823	823	1448	2408	2409	3353

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	Top Hit Descriptor	Homo sapiens partial 5-HT4 receptor gene, exons 2 to 5	Homo sepiens solute carrier family 13 (sodium-dependent dicerboxylate transporter), member 2 (SLC13A2) mRNA	Homo sepiens ubiquitin specific protease 9, X chromosome (Drosophila fat facets related) (USP9X), mRNA	Homo sapiens phosphoribosy pyrophosphate synthetase 2 (PRPS2), mRNA	Homo sapiens eukaryotic translation initiation factor 2B, subunit 2 (beta, 39kD) (EIF2B2), mRNA	Homo sapiens eukaryotic translation Initiation factor 2B, subunit 2 (beta, 39kD) (EIF2B2), mRNA	Homo sapiens mRNA for KIAA1263 protein, partial cds	H.sapiens flow-sorted chromosome 6 Hindill fragment, SC6pA16D3	H.sapiens flow-sorted chromosome 5 Hindill fragment, SO6pA16D3	Homo sapiens putative nuclear protein (HRIHFB2122), mRNA	Homo sapiens non-histone chromosome protein 2 (S. cerevisiae)-like 1 (NHP2L1), mRNA	Homo sapiens cadherin EGF LAG seven-pass G-type receptor 1 (CELSR1), mRNA	Homo sapiens calcineurin binding protein 1 (KIAA0330), mRNA	Homo sapiens calcineurin binding protein 1 (KIAA0330), mRNA	Homo capiens low density lipoprotein-related protein 2 (LRP2), mRNA	Homo sapiens neurofibromin 2 (bilateral acoustic neuroma) (NF2) mRNA	Homo sapiens mRNA for KIAA1476 protein, peirtial cds	Homo sapiens mRNA for KIAA1476 protein, pertial cds	Human cyclophilin-related processed pseudogene	wa33104.x1 NCI_CGAP_Kld11 Homo saplens CDNA clone IMAGE:2299903 3' similar to contains THR.t2	I HK repetitive element ;	Homo sapiens chromosome 21 segment HS21C084	RC0-BN0284-300500-031-e05 BN0284 Homo sapiens cDNA	RC0-BN0284-300500-031-e05 BN0284 Homo sapiens cDNA	Homo sapiens mannosidase, beta A, tysosomai (MANBA) gene, and ubiquitin-conjugating enzyme E2D 3	(UBE2D3) genes, complete cds	QV4-BT0257-081199-017-e03 BT0257 Homo saplens cDNA	Homo sapiens intersectin 2 (SH3D1B) mRNA, complete cds	Homo sapiens ADP/ATP carrier protein (ANT-2) gene, complete cds	af70e11.r1 Scares_NhHMPu_S1 Homo sapienis cDNA clone IMAGE:1047404 6' similar to WP:K01H12.1 CE03453:	EST_HUMAN DKFZp566F104_r1 568 (synonym: hfkd2) Homp saplens cDNA clone DKFZp566F104 5'
	Top Hit Database Source	NT	ΤN	LN	LN	NT	IN	IN	NT	NT.	NT	LN TN	NT	NT	NT	N	TN	ΙN	N	Ν		EST_HUMAN	N	EST_HUMAN	EST_HUMAN		IN	EST_HUMAN	NT	N	EST HUMAN	EST_HUMAN
	Top Hit Acession No.	AJ243213.1	4506978 NT	11420654 NT	11421041 NT	7657057 NT	7657057 NT	AB033089.1	Z78766.1	Z78766.1	11418086 NT	11418192 NT	11418322 NT	11417862 NT	11417862 NT	11430460 NT	4557794 NT	AB040909.1	AB040909.1	X52858.1		AI632733.1	AL163284.2	BF329911.1	BF329911.1		AF224669.1	BF330676.1	AF248540.1	L78810.1	AA625207.1	AL039044.1
Ö	Most Similar (Top) Hit BLAST E Value	4.0E-62	4.0E-62	4.0E-62	4.0E-62	4.0E-62	4.0E-62	4.0E-82		4.0E-62	4.0E-82	4.0E-62	4.0E-62	4.0E-62	4.0E-62	4.0E-62	3.0E-62	3.0E-62		3.0E-62		_	_		2.0E-62				1.0E-62	1.0E-62	1.0E-62	
	Expression Signal	2.03	1.66	2.42	1.68	2.21	2.21	6.3	2.43	2.43	2.95	2.98	1.99	4.2	4.2	1.61	1.12	98.0	0.95	1.92		4.35	1.5	4.8	4.8		3.94	8.93	1.24	6.83	1.02	1.12
	ORF SEQ ID NO:		25608	25822	26273	26528	26529	27204	28505	28506	28792		25201	25198	25189	25210	19868	22728	22729	23340		27038	20969	27165	27168				20791	21288	21628	22606
	Exan SEQ ID NO:	14824	15524	15709	16120	16357	16357	17011	18254	18254	18891	19578	19305	19302	19302	19335	10053	12936	12936	13554	-	16846	118	16973	16973		17696	18744	10946	11431	11667	12811
	Probe SEQ ID NO:	4947	5609	5804	6254	6498	6498	7134	8377	8377	9135	8322	8743	9792	9792	9835	89	3008	3008	3640	0000	200	12	9602	7096		7846	8936	1028	1526	1768	2884

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			Γ	Γ	Γ		Γ	Γ		Γ			Γ			Γ	Γ	Γ	Ī	Γ	İ	Γ	Γ	Ī	Γ			Ţ	Τ	T	Γ	Γ	Γ
	Top Hit Descriptor	Homo sapiens hypothetical protein FLJ20212 (FLJ20212), mRNA	zg89f10.s1 Soares_fetal_heart_NbHH19W Homo sepiens cDNA clone IMAGE:409771 3'	2089/10.s1 Soares_fetal_heart_NbHH19W Homo septens cDNA clone IMAGE:409771 3'	Homo saplens KIAA0763 gene product (KIAA0763), mRNA	Homo sapiens KIAA0763 gene product (KIAA0763), mRNA	H.saplens lysosomal acid phosphatase gene (EC 3.1.3.2) Exon 9	H.sapiens lycosomal acid phosphatase gene (EC 3.1.3.2) Exon 9	aa33d08.51 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:815055 3'	H.sapiens flow-sorted chromosome 6 Hindill fragment, SC6pA14D8	Homo sapiens cadherin EGF LAG seven-pass G-type receptor 1 (CELSR1), mRNA	Homo sapiens law density lipopratein-related protein 2 (LRP2), mRNA	QV4-ST0234-181199-037-f05 ST0234 Homo sapiens cDNA	C18159 Human placenta cDNA (TFujiwara) Homo sapiens cDNA clone GEN-558C10 5'	Homo sapiens mRNA for KIAA0350 protein, partial cds	Homo saplens mRNA for KIAA0350 protein, partial cds	Homo sapiens aconitase 2, mitochondrial (ACC2), mRNA	Homo sapiens mRNA for PkB kinase	Homo saplens nucleoporin 88kD (NUP88), mRNA	Homo saplens Ras association (RaiGDS/AF-6) domain family 2 (RASSF2), mRNA	Homo sapiens monoamine oxidase A (MAOA), nuclear gene encoding mitochondrial protein, mRNA	Homo sapiens IL2-Inducible T-cell kinase (ITK) mRNA	Gallus gallus Dach2 protein (Dach2) mRNA, complete cds	Gallus gallus Dach2 protein (Dach2) mRNA, ccimplete cds	Homo saplens chromosome 21 segment HS21(2068	wm55g11.x1 NCL_CGAP_Ut2 Homo sapiens cIDNA clone IMAGE:2439908 3'	ne83f02.r1 NCI_CGAP_Pr1 Homo sapiens cDNA clone IMAGE:745947 similar to gb: Y00361 60S RIBOSOMAL PROTEIN (HUMAN):	Homo saplens chromosome 21 segment HS210078	CM3-BT0595-190100-072-a09 BT0595 Hamo sapiens cDNA	CM3-BT0595-190100-072-a09 BT0595 Homo sapiens cDNA	UI-H-BI1-abq-a-02-Q-UI.s1 NCI_CGAP_Sub3 Homo sapiens cDNA clone IMAGE:2712482 3'	UI-H-BI1-abq-a-02-0-UI.s1 NCI_CGAP_Sub3 Homo sapiens cDNA clone IMAGE:2712482 3'	Homo saplens mRNA for KIAA0717 protein, paritial cds
פון באטון ו	Top Hit Database Source	NT	EST_HUMAN	EST_HUMAN	NT	TN	NT	LN LN	EST_HUMAN	TN	LY	TN.	EST_HUMAN	EST_HUMAN	NT	NT	FZ	N F	FZ	TN	Ā	N P	TN.	TN	IN	EST_HUMAN	NAMINE TOTAL	LN	EST HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	L
5	Top Hit Aœssion No.	8923201 NT	AA722878.1	AA722878.1	7662289 NT	7662289 NT	X15533.1	X16533.1	AA465170.1	Z78698.1	11418322 NT	11430460 NT	AW816405.1	C18159.1	AB002348.2	AB002348.2	1418185	Y15056.1	11426985 NT	11421160 NT	4557734 NT	5031810 NT	AF198349.1		AL163268.2	AI872137.1	A A 4 2 0 8 0 3 4	Ι	Ĺ	AW 750372.1	AW134709.1	AW134709.1	AB018280.1
•	Most Similar (Top) Hit BLAST E Value	1.0E-62	1.0E-62	1.0E-62	1.0E-62	1.0E-62	1.0E-62)	1.0E-62	1.0E-62		1.0E-62	1.0E-82	9.0E-63	9.0E-63	9.0E-63	9.0E-83		9.0E-83	9.0E-63	9.0E-63	8.0E-63	8.0E-63				7.0E-63	8.0F-63.4	_			-	4.0E-83	3.0E-63 A
	Expression Signal	1.32	2.17	2.17	1.53	1.53	1.81	1.81	2.81	2.13	1.94	2.3	2.14	1.51	7.42	7.42	6.51	1.31	4.39	1.37	1.52	2.17	3.81	3.81	3.37	1.78	40.61	86.0	2.88	2.88	2	2	1.97
	ORF SEQ ID NO:		26251	26252		27311		27332		28845					23638	23639		25104	26281	26917	22077	22102		23135				23001					21668
	Exan SEQ ID NO:	14321	16102	16102	17116	17116	17139	17139	17300	18561	19199	18322	10294	12179	13863	13863	15088	15274	16127	16724	12178	12203	13332	13332	14068	10837	15196	13200	15785	15785	18347	18347	11788
	Probe SEQ ID NO:	4426	9539	6236	7238	7239	7282	7262	7512	8673	8823	9815	335	2297	3955	3826	6210	5354	6262	6845	2296	2322	3416	3416	4168	913	5274	3279	5879	5879	8474	8474	1893

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Top Hit Descriptor	Human Met-fRNA-i gene 1	Homo sapiens zinc finger protein 144 (Mel-18) (ZNF144), mRNA	Homo sapians hapatocellular carcinoma antigen gene 520 (LOC63928), mRNA	601485656F1 NIH_MGC_69 Homo sepiens ¢DNA clone IMAGE:3888253 5'	601485656F1 NIH_MGC_69 Homo saplens cDNA clone IMAGE:3888253 5'	Human DNA topolsomerase I mRNA, partial cds	Homo sapiens eyes absent (Drosophila) homolog 2 (EYA2), mRNA	Homo sapiens glutamate-cysteine ligase (garrima-glutamylcysteine synthetase), catalytic (72,8kD) (GLCLC) mRNA	Homo sapiens Down syndrome candidate region 1 (DSCR1), mRNA	Homo sapiens RHCE mRNA for Rh blood CE group antigen polypeptide, complete cds	Homo saplens RHCE mRNA for Rh blood CE group antigen polypeptide, complete cds	Homo sapiens emyold beta (A4) precursor protein (protease nexin-II, Alzheimer disease) (APP), mRNA	Homo sapiens chromosome 3 subtelomeric region	Homo sapiens polycystic kldney disease-associated protein (PKD1) gene, complete cds	Homo sapiens Jun dimerization protein gene, partial cds; cfos gene, complete cds; and unknown gene	QV1-FT0170-040700-265-c05 FT0170 Homo sapiens cDNA	QV1-FT0170-040700-265-c05 FT0170 Homo sapiens cDNA	Human germline T-cell receptor beta chain Dopamine-beta-hydroxylase-like, TRY1, TRY2, TRY3,	TCRBVZ/S1P, 1CKBVZ2S1AZN11, 1CKBVSS1A11, 1CKBV/S1A1NZ1, 1CKBVSS1A11, 1CKBV 1353, 1 TCRBV6S7P, TCRBV7S3A2T, TCRBV13S2A1T, TCRBV9S2A2PT, TCRBV7S2A1N4T,	TCRBV13S9/13S>	Homo sapiens Carbonic anhydrase-related protein 10 (LOC56934), mRNA	Homo sapiens Carbonic anhydrase-related protein 10 (LOC56934), mRNA	Homo sapiens chromosome 21 segment HS21,0010	zb18b05.s1 Soares_fetal_lung_NbHL19W Homo sapiens cDNA clone IMAGE:302385 3' sImilar to	gb:X17206 40S RIBOSOMAL PROTEIN S4 (HUMAN);	Homo sapiens neurexin III-alpha gene, partial cds	Homo sapiens neurexin III-alpha gene, partial cds	Homo sapiens aconitase 2, mitochondrial (ACO2), mRNA	Homo sapiens calcium channel, voltege-dependent, alpha 11 subunit (CACNA11), mKNA	HSCZVD111 normalized Infant brain cDNA Homo sapiens cDNA clone c-zvd11
Top Hit Database Source	Į.				EST_HUMAN	NT					FX		NT	TN	TN					뉟	F	۲	Z	, T	T HUMAN	Ę	M	Ā		EST_HUMAN
Top Hit Acession No.	J00310.1	6005963 NT	11545810 NT			2.0E-63 U07804.1	4885226 NT	4557624 NT	7657042 NT	2.0E-63 AB030388.1	2.0E-63 AB030388.1	4502166 NT	4F109718.1	39891.1	2.0E-63 AF111167.2	3F373541.1	2.0E-63 BF373541.1			J66059.1	9910365 NT	9910365 NT	2.0E-63 AL163210.2		2.0E-63 N78945.1	2.0E-63 AF099810.1	2.0E-63 AF099810.1	11418185 NT	11418157 NT	1.0E-63 F08485.1
Most Similar (Top) Hit BLAST E Value	3.0E-63	3.0E-83	3.0E-63	3.0E-63	3.0E-63	2.0E-63	2.0E-83	2.0E-63	2.0E-63	2.0E-63	2.0E-63 /	2.0E-63	2.0E-63 A	2.0E-63 L39891.1	2.0E-63	2.0E-63	2.0E-63			2.0E-63	2.0E-63	2.0E-63	2.0E-63		2.0E-63					
Expression Signal	1,34	8.97	27.69	1.82	1.82	1.11	1.74	1.39	528	2.47	2.47	1.68	1.78	1.86	1.13	2.45	2.45			1.37	1.41	1.41	3.8		12.54	3.02	3.02	10.85	1.39	291
ORF SEQ ID NO:		20978	25928	27683	27664	19972					21313		22970	L		25578				26061	L		27034			28292	28293	25058		Ц
Exan SEQ ID NO:	12615	L	Ι.	17448	17448	<u>L</u>	L	10432		L		1	1	ĺ	<u> </u>	15502	L	L		15929		İ.	<u> </u>	<u> </u>	18017	18042	18042	L		Ш
Probe SEQ ID NO:	2753	2791	5896	7697	7697	184	191	489	808	1647	1647	3118	3248	3831	4760	5587	5587			6025	6208	6208	6964		8129	8154	8154	9243	9864	4246

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Top Hit Descriptor	HSCZVD111 normalized infant brain cDNA Homo sapiens cDNA clone c-zvd11	QV0-ST0215-060100-083-b09 ST0215 Hamp sapiens cDNA	Homo sapiens chromosome 21 segment HS21C047	Homo sapiens chromosome 21 segment HS21C007	tm50b07x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2161525 3'	601155232F1 NIH_MGC_21 Homo saplens cDNA clone IMAGE:3139038 5'	601508968F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3910336 5	Homo sapiens Ran GTPase activating protein 1 (RANGAP1), mRNA	yb98b02.r1 Stratagene lung (#937210) Homo sapiens cDNA clone IMAGE:79179 5'	601311455F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3633204 5	Homo sapiens thimet oligopeptidase 1 (THOP1) mRNA	Homo sapiens thimet oligopeptidase 1 (THOP1) mRNA	Homo sapiens EWS, gar22, rrp22 and bam22 genes	wb51e07.x1 NCI_CGAP_GC6 Homo septems cDNA clone IMAGE:2309220 3' similar to gb:M15182 BETA-	URUNIDASE PRECURSOR (HUMAN);	wb51e07.x1 NCI_CGAP_GC6 Homo sepiens cDNA clone IMAGE:2309220 3' similar to gb:M15182 BETA-GLUCURONIDASE PRECURSOR (HUMAN);	wv13e03.x1 NCI_CGAP_Brn23 Homo sapiens cDNA clone IMAGE:2529436 3'	wv13e03.x1 NCI_CGAP_Brn23 Homo saplens cDNA clone IMAGE:2529436 3'	Homo sapiens MCP-1 gene and enhancer region	Homo sapiens MCP-1 gene and enhancer región	Homo sapians protein kinase C beta-II type (PRKCB1) mRNA, complete cds	Homo sapiens mesenchyme homeo box 1 (MEOX1), mRNA	Homo sapiens mesenchyme homeo box 1 (MEOX1), mRNA	Homo saplens acetyl-CoA synthetase (LOC55902), mRNA	Homo saplens progressive ankylosis-like protein (ANK) mRNA, complete cds	trkC [human, brain, mRNA, 2715 nt]	Homo sapiens stromal antigen 3 (STAG3), mRNA	Homo sapiens stromal antigen 3 (STAG3), mRNA	Homo sapiens interleukin 10 receptor, beta (IL19RB), mRNA	Homo sapiens chromosome 21 unknown mRNĄ	Hamo saplens chramosame 21 unknown mRNA	Homo saplens phosphoglucomutase-related protein (PGMRP) gene, complete cds	Hamo saplens phosphoglucomutase-related protein (PGMRP) gene, complete cds
± 9 0	Τ		Homo sapiens chromoson	Homo sapiens chromoson	Г					Г	Г	Homo sapiens thimet oligo	Homo sapiens EWS, gar2	Г	7		Г		Homo sapiens MCP-1 ger	Homo sapiens MCP-1 ger	Homo sapiens protein kina	Homo sapiens mesenchyr	Homo sapiens mesenchyr	Homo saplens acetyl-CoA	Homo saplens progressive	trkC [human, brain, mRN/	Homo sapiens stromal ant	Homo sapiens stromal ant	Homo sapiens interleukin	Homo sapiens chromoson	Homo sapiens chromoson	Homo saplens phosphoglu	Homo sapiens phosphogia
Top Hit Database Source	EST_HUMAN	EST_HUMAN	ΝΤ	NT	EST_HUMAN	EST_HUM	EST_HUMAN	'NT	EST_HUMAN	EST_HUMAN	N.	NT.	NT TN	100	ESI TOWAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	Ρ	LΝ	NT	INT	NT	NT	NT	NT	N.	INT	NT	NT	NT	N	N
Top Hit Acession No.	F08485.1	AW582266.1	AL163247.2	AL163207.2	AI478186.1	ш	BE885755.1	11418177 NT	T60651.1	BE394321.1	4507490	4507490 NT	Y07848.1	A 1054000 4	AI651892.1	Al651992.1	AW026445.1	AW026445.1	Y18933.1	Y18933.1	M13975.1	11525879 NT	11525879 NT	11420555 NT	AF274753.1	S76475.1		11420197 NT	11526198 NT	AF231919.1	AF231919.1	L40933.1	L40933.1
Most Similar (Top) Hit BLAST E Value	1.0E-63	1.0E-63	1.0E-63	1.0E-83	9.0E-64	8.0E-64	8.0E-64	8.0E-64	8.0E-64	7.0E-64	7.0E-64	7.0E-84	7.0E-64	70 10 0	6.0E-64	6.0E-64	6.0E-64	6.0E-64	6.0E-84	6.0E-64	6.0E-64	6.0E-64	6.0E-64	6.0E-84	6.0E-64	6.0E-64	6.0E-64	6.0E-64	6.0E-64	5.0E-64	5.0E-64	5.0E-64	5.0E-64
Expression Signal	2.91	1.39	2.3	3.02	4.78	7.89	3.16	2.61	2.58	0.99	2.44	2.44	2.13		-	1.7	3.7	3.7	2.64	2.64	14.41	2.58	2.58	7.8	2.08	2.34	7.57	7.57	4.08	2.44	2.44	2.42	2.42
ORF SEQ ID NO:		25509			26681		25747					54304			21463	21464	12801			75434	25444	26312	26313			27669	28287	28288	25321	20574		21167	21168
Exan SEQ ID NO:	L	15443	16813	19633	16471	10948	15641	18841	18875	13402	14613	14513	17616	L	1384	11594	13011				15384		16157					18039	18967				11307
Probe SEQ ID NO:	4248	5526	6935	9879	6591	1030	5733	8068	9111	3486	4625	4625	7766	1600	7801	1692	3084	3084	5454	5454	5464	6283	6283	7376	7472	7604	8151	8151	9262	803	803	1402	1402

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	Top Hit Descriptor	Human ((3)mbt protein homolog mRNA, complete cds	Homo sapiens KIAA0618 gene product (KIAA0618), mRNA	Homo sapiens KIAA0618 gene product (KIAA0618), mRNA	Homo sapiens putative transcription factor CR53 (CR53) mRNA, partial cds	RC3-ST0197-120200-015-a03 ST0197 Homp sapiens cDNA	RC3-ST0197-120200-015-a03 ST0197 Homp saplens cDNA	C18895 Human placenta cDNA (TFujiwara) Homo sapiens cDNA clone GEN-569E02 5	601589565F1 NIH_MGC_7 Homo sapiens dDNA clone IMAGE:3943577 5'	AV711714 DCA Homo sapiens cDNA clone DCAAMO01 5'	AV711714 DCA Homo sapiens cDNA clone DCAAMO01 5'	H.sapiens isoform 1 gene for L-type calcium channel, exon 28	RC6-FN0019-290600-011-G11 FN0019 Homo saplens cDNA	Homo sapiens golgi matrix protein GM130 (GOLGA2) mRNA, complete cds	Homo sapiens golgi matrix protein GM130 (GOLGA2) mRNA, complete cds	bb/2h12.y1 NIH_MGC_12 Homo sepiens cDNA clone IMAGE:3047975 5' similar to gb:L08069 DNAJ	NOTEIN TOWING & (TOWAN),	bb72h12.y1 NiH_MGC_12 Homo sapiens cDNA clone IMAGE:3047975 5' similar to gb:L08069 DNAJ PROTEIN HOMOLOG 2 (HUMAN):	Homo sablens chromosome 21 segment HS21C046	Homo saplens chromosome 21 segment HS2; C046	Hamo sapiens chromosame 21 segment HS21 C048	Hamo sapiens chromosome 21 segment HS21C046	Homo sapiens chromosome 21 segment HS21C027	af09d08.s1 Soares_testis_NHT Homo saplens cDNA clone IMAGE:1031151 3'	Homo saplens elF4E-like cap-binding protein (4EHP) mRNA	wo87b01.x1 NCI_CGAP_Kid11 Homo sapiens oDNA clone IMAGE:2462281 3' similar to contains element	L1 repetitive element;	Homo sapiens chromosome 21 segment HS21,0048	Homo sapiens chromosome 21 segment HS21,0046	Homo sapiens glutamic-oxaloacetic transaminase 2, mitochondrial (aspartate aminotransferase 2) (GOT2), nuclear gene encoding mitochondrial protein, mRNA	EST370215 MAGE resequences, MAGE Homo sapiens cDNA	EST370216 MAGE resequences, MAGE Hamo sepiens cDNA	AU124387 NT2RM2 Homo saplens cDNA clone NT2RM2002113 5'
	Top Hit Database Source	E				EST_HUMAN F	Г	г			EST_HUMAN /	NT I	EST_HUMAN F		L L	TOT LINAMI	┪	EST HUMAN	T			±N	NT IN	EST_HUMAN			T_HUMAN		NT		T HUMAN	П	EST_HUMAN /
3	Top Hit Acession No.	U89358.1	7662205 NT	7882205 NT	AF017433.1	AW813783.1	AW813783.1	C18895.1	BE794381.1	AV711714.1	AV711714.1	Z26273.1	7.	AF248953.1	AF248953.1	0.000004	5EZ005Z1.1	BF208521 1				AL163246.2	AL163227.2	AA609940.1	4757701 NT				AL163246.2	4504068 NT	AW958145.1		AU124387.1
	Most Similar (Top) Hit BLAST E Value	5.0E-64	5.0E-84	5.0E-64		4.0E-64	4.0E-64	3.0E-64	3.0E-64	3.0E-64				3.0E-64			-	3 0F-64	-	3.0E-64	3.0E-84	3.0E-64	3.0E-64	2.0E-64	2.0E-64		2.0E-64		20E-64	2.0E-64	2.0E-64	2.0E-64	20E-64
	Expression Signal	1.67	2.68	2.66	5.61	3.9	3.9	3.85	0.95	1.51	1.51	1.34	3.39	1.81	1.81		2		1 28	1.26	1.76	1.76	4.59	0.94	1.32		1.78	3.03	3.03	2.56	1.33	1.33	2.62
	ORF SEQ ID NO:	21457	21235	21236	23575	28324	28325	21836	22943	23112	23113	25713	25942	27005	27006	97050	2/010	27017			28703	28704	29040	20831	21137			25252	22253	22823	23416		
	Exan SEQ ID NO:	11585	11370	11370	13787		ı	12039	13140			1			16811	i	10024	16824	17280	17289	18434	18434	18746	10988	11282		12354	12359	12359	13027		H	15565
	Probe SEQ ID NO:	1683	2796	2798	3876	8188	8188	2151	3216	3396	3396	5703	5911	6933	8833	8748	Op/RO	6948	7422	7422	8565	8565	8638	1072	1376		2478	2484	2484	3101	3718	3719	5653

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Table 4
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RF SEQ Expression (Top) Hit Acession (Top) Hit Acession Signal BLASTE No. Signal Average Source Source	26799 1.3 2.0E-64 AF113708.1 NT Homo sepiens engiopoletin 4 (ANG4) mRNA, partial cds	4.97 2.0E-64	1.31	26060 3.88 2.0E-84 M77185.1 NT H.sapiens dopamine receptor D5 pseudogene 1, partial cds	BF528114.1 EST_HUMAN	6.4 2.0E-64 AI922911.1 EST_HUMAN	6.4 2.0E-64 Al922911	25347 1.73 2.0E-64 8567387 Homo saplens period (Drosophila) homolog 3 (PER3), mRNA			21508 5.88 1.0E-64 Al929419.1 EST_HUMAN gb:L21696_cds1 PROTHYMOSIN ALPHA (HUMAN);contains element MSR1 repetitive element;	Homo sapiens transcription factor IGHM enhancer 3, JM11 protein, JM5 protein, T64 protein, JM10 protein, A4 differentiation-denandent protein, title I M domain protein 8, and expensive sense.	23188 5.61 1.0E-64 AF188779.1 INT complete cds; and L-type calcium channel a>	23248 1.32 1.0E-64 AF228627.1 NT Homo sapiens TRIAD3 mRNA, partial cds		1.62 1.0E-64 AL163246.2 NT	0.93 9.0E-65 X89211.1	22018 0.93 9.0E-65 X89211.1 (NT H.sapiens DNA for endogenous retroviral like element	15.1 9.0E-45 BF330676.1 EST_HUMAN QV4-BT0257-081199-017-e03 BT0257 Homo saplens cDNA	28903 10.83 8.0E-65 AI929244.1 EST_HUMAN SW:RL21_HUMAN P46778 60S RIBOSOMAL PROTEIN L21.;	27936 2.05 7.0E-65 BE081653.1 EST_HUMAN QV2-BT0635-240400-162-602 BT0635 Homo saplens cDNA	1.52 6.0E-65 AV721898.1	R.32 6.0E-65 AA550929.1 EST_HUMAN RIBOSOMAL PROTEIN L32 (HUMAN);	xc07b09.x1 NCI_CGAP_Co21 Homo septens cDNA clone IMACE:2583545 3' similar to TR:Q63306 Q63306 2559 6.0E-65 AW083252.1 EST HUMAN LONG INTERSPERSED REPETITIVE DNA CONTAINING 7 ORF'S :contains L1.b2.L1 repetitive element:	4.25 6.0E-65 AA427878.1 EST_HUMAN	4.25 6.0E-65 AA427878.1 EST_HUMAN	A 18 A DE AS BESTA18 1 EST MINAN
ORF SEQ Express ID NO: Sign	25799	25938	26004	26060	28279	28534	28635	25347		20039	21508		23188	23248	23249		22017	22018						27150	27275	27276	28375
Probe Exan OF SEQ ID NO:	5783 15689		5976 15880	6024 15928	8144 18032		8406 18282	9182 18921	9617 19195	268 10224	1740 11641		3466 13382		3536 13452				8822 18635	8789 18613	7841 17891	1040 10958	1880 11776	7080 16957	7209 17086	7209 17086	12427 18437

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Ì) 		
Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
615	10551	20362		5.0E-65	AF064604.1	NT	Homo sapiens KE03 protain mRNA, partial cds
1331	11238	21094	1.62	5.0E-65	7861951 NT	NT	Homo sapiens KIAA0158 gene product (KIAA0156), mRNA
1331	11238	21085		5.0E-65	7881951 NT	NT	Hamo saplens KIAA0156 gene product (KIAA0156), mRNA
2110	11999	21898	1.02	5.0E-65	AB033768.1	NT	Homo saplens hPAD-colony10 mRNA for peptidylarginine deiminase type I, complete cds
3217	13141		1.91	5.0E-65		NT	Homo saplens ubiquitin specific protease 13 (Isopeptidase T-3) (USP13) mRNA
3217	13141	L	1.91	5.0E-65	4507848 NT	NT	Homo sapiens ubiquitin specific protease 13 (Isopeptidase T-3) (USP13) mRNA
186	10158		1.09	4.0E-65	AL120419.1	EST_HUMAN	DKFZp761G108_r1 761 (synonym: hamy2) Homo sapiens cDNA clone DKFZp761G108 5'
728	10660	20491	1.58		4.0E-65 AI268468.1	EST_HUMAN	qm46e01.xf Sogres_placenta_8to9weeks_2t\bHP8to9W Homo sapiens cDNA clone IMAGE:1891800 3'
728	10860	20492	1.56		AI268468.1	EST_HUMAN	qm46e01.x1 Soeres_placenta_8to9weeks_2NbHP8tb9W Homo sapiens cDNA clone IMAGE:1891800 3'
1082	10978	20822	1.88		4826735 NT	LN	Homo sapiens fragile X mental retardation, autosomal homolog 1 (FXR1), mRNA
1470	11375	21240		4.0E-65	4506636 NT	NT	Homo sapiens ribosomal protein L34 (RPL34) mRNA
2288	12171	22068	1.03	4.0E-65	BE221469.1	EST_HUMAN	hu25e04.x1 NCI_CGAP_Me115 Homo sapiens cDNA clone IMAGE:3171102 31
2288	12171				3E22146	EST_HUMAN	hu25e04.x1 NCI_CGAP_Mel15 Homo sapien's cDNA clone IMAGE:3171102 3'
5138	15005		0.95	4.0E-65	9055269 NT	NT	Homo sapiens low density lipoprotein receptor related protein-deleted in tumor (LRPDIT), mRNA
5138	15005				, 9055269	니	Homo sapiens low density lipoprotein receptor related protein-deleted in tumor (LRPDIT), mRNA
5740	15648		3.93	4.0E-65	AB033093.1	NT	Homo sapiens mRNA for KIAA1267 protein, partial cds
5740	15648	25755	3.93	4.0E-65	AB0330	NT	Homo sapiens mRNA for KIAA1267 protein, partial cds
6285	16149	26304	2.28		11545780 NT	NT	Homo sapiens hypothetical protein FLJ22087 (FLJ22087), mRNA
8020	17870		2.17	4.0E-65	AJ277546.2	NT	Homo sapiens WEE1 gene for protein kinase and partial ZNF143 gene for zinc finger transcription factor
8446	18320		7.47	4.0E-65	AF119846.1	IN	Homo sapiens PRO1474 mRNA, complete cds
9471	10978	20822	1.34	4.0E-85	4826735 NT	NT	Homo sapiens fragile X mental retardation, autosomal homolog 1 (FXR1), mRNA
8975	19434				11430460 NT	NT	Homo sapiens low density lipoprotein-related protein 2 (LRP2), mRNA
1212	12646		6.37	3.0E-65	X78932.1	TN	H.saplens HZF9 mRNA for zinc finger protein
1780	11679	21557	1.14	3.0E-65	A1000692.1	EST HUMAN	ov23f03.s1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1638173 3' similar to contains element MSR1 repetitive element ;
3239	13162	<u> </u>	1.39		4504950 NT	NT	Homo sapiens laminin, beta 1 (LAMB1), mRNA
3660	13574		0.98		A1000692.1	EST_HUMAN	ov23f03.s1 Soares_testis_NHT Homo eaplens cDNA clone IMAGE:1638173 3' similar to contains element MSR1 repetitive element ;
4551	14444		1.41		TN 6912385 NT	NT	Homo sapiens rab6 GTPase activating protein (GAP and centrosome-associated) (GAPCENA), mRNA
7783	17843	27876		3.0E-65	3E78736	EST_HUMAN	601479686F1 NIH_MGC_68 Homo sapiens ciDNA clone IMAGE:3882405 5'

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Γ		П	\neg						Т				Γ	Γ	Γ	П			П				٦		٦			0	T	Т	T	Т	٦
	Top Hit Descriptor	zw65a06.r1 Soares_testis_NHT Homo seplens cDNA clone IMAGE:781042 6	602/155062F1 NIH_MGC_83 Homo saplens cDNA clone IMAGE:4295968 5'	601190883F1 NIH MGC_7 Homo saplens clDNA clone IMAGE:3634741 5'	602134359F1 NIH_MGC_81 Homo saplens cDNA clone IMAGE:4289295 6'	Homo sapiens mRNA for FLJ00056 protein, partial cds	Homo sapiens mRNA for FLJ00056 protein, partial cds	Homo sapiens SWI/SNF related, matrix associated, actin dependent regulator of chromatin, subfamily d,	member 3 (SMARCD3), mRNA	EST178755 Colon carcinoma (HCC) cell line Homo saplens cDNA 5' end similar to similar to endogenous retrovinis	601854033F1 NIH MGC 57 Homo sapiens cDNA done IMAGE:4073769 5'	1601763488F1 NIH MGC 20 Homo septens CDNA clone IMAGE: 4028501 5	Homo sapiens putative Rab5 GDP/GTP exchange factor homologue (RABEX5), mRNA	Homo sapiens mRNA for KIAA1513 protein, partial cds	hz24a09.x1 NCI_CGAP_GC6 Homo saplens cDNA clone IMAGE:3208888 3'	Homo sapiens glypican 4 (GPC4) mRNA	Homo sapiens glypican 4 (GPC4) mRNA	wx09c09 x1 NCI_CGAP_Gas4 Homo sapiens cDNA clone IMAGE:2543152 3'	wx09c09.x1 NCI_CGAP_Gas4 Homo saplens cDNA clone IMAGE:2543152 3'	QV2-ST0298-140200-042-f12 ST0298 Homo saplens cDNA	QV2-ST0298-140200-042-f12 ST0298 Homo saplens cDNA	AU141295 THYRO1 Homo saplens cDNA clone THYRO1000356 5	AU141295 THYRO1 Homo saplens cDNA clone THYRO1000356 5'	602126239F1 NIH_MGC_56 Homo saplens cDNA clone IMAGE:4283313 5'	AU128040 NT2RP2 Homo sapiens cDNA clone NT2RP2004714 5	AU129040 NT2RP2 Homo sapiens cDNA clone NT2RP2004714 5'	Homo sapiens inositol 1,4,5-triphosphate receptor, type 1 (ITPR1), mRNA	dd56a02.x1 Soares_testis_NHT Home sapiens cDNA clone IMAGE:1733450 3' similar to gb:M29581 ZINC	FINGER PROJEIN 8 (HOWAN), contents MERIES I MERIES IN FEBRUARE GETTERS.	AU153793 NT2RP3 Homo seplens cDNA clorie NT2RP3004016 3'	Human platelet factor 4 varation 1 (PF4var1) gene, complete cds	Homo sapiens ribosomal protein L7a (RPL7A) mRNA	602126239F1 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:4283313 5
	Top Hit Database Source	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	N	LZ LZ		LN.	NAM II	EST HUMAN	EST HIMAN	LN	LN LN	EST HUMAN	LN	N	EST HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	닐		EST_HUMAN	EST_HUMAN	TN	INT	EST HUMAN
	Top Hit Acession No.	AA430006.1	BF680294.1	BE263373.1	BF576922.1	AK024463.1	AK024463.1		11419247 NT	A A 207004 4	BF246086 1	RE105544 1	7657495 NT	AB040946.1	BE466681.1	4504082 NT	4504082 NT	AW028340.1	AW029340.1	AW820481.1	AW820481.1	AU141295.1	AU141295.1	BF698707.1	AU129040.1	AU129040.1	11431994 NT		AI191716.1	AU153793.1	M26167.1	4506660 NT	BF698707.1
	Most Similar (Top) Hit BLAST E Value	3.0E-65	2.0E-65	2.0E-65	2.0E-65	2.0E-65	2.0E-65		2.0E-85	2000							1.0E-65		1.0E-85			1.0E-65	1.0E-65	1.0E-65	1.0E-65	1.0E-65	1.0E-65		1.0E-65		1.0E-65		1.0E-85
	Expression Signal	9.69	5.27	4.55	23.12	1.27	127		2.85	4	237	80,	1.32	0.91	0.89	2.13	2.13	2.37	2.37	1.58	1.68	2.38	2.38	1.76	1.62	1.62	2.83		9	1.25	2.23	12.89	2.18
	ORF SEQ ID NO:	28119	23078		26247				28189				20282						L		L	L	26915	27201	27282				ļ		28181		28610
	Exan SEQ ID NO:	17877	13277	16843		}_	17010	1	17939	40074	10504	40080	10470	11888	1	ı	13826	I	1_	L	16681	١.		17009	17092	17092	17099	1	1	17537	17933	18046	18346
	Probe SEQ ID NO:	8693	3368	5938	6233	7133	7133		8048	2000	o San	1	578	1804	3326	3917	3917	4112	4112	6802	6802	6842	6842	7132	7215	7215	7222		7456	7687	8042	8158	8473

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П			Γ	Г	Γ	Π						Γ	Г	Г		\neg		Τ		Π		П							П			T
	Top Hit Descriptor	ts76a08.x1 NCI_CGAP_GC6 Homo sepiens cDNA clone IMAGE:2237170 3' similar to gb:L15533_ma1 PANCREATITIS ASSOCIATED PROTEIN 1 PRECURSOR (HUMAN);	Homo sapiens TNF-Inducible protein CG12-1 (CG12-1), mRNA	Homo sapiens cadherin EGF LAG seven-pass G-type receptor 1 (CELSR1), mRNA	Homo sapiens sulfotransferase-related protein (SULTX3), mRNA	Novel human gene mapping to chomosome 22	Novel human gene mapping to chomosome 22	Homo sapiens 26S proteasome-associated ped1 homolog (POH1) mRNA	Homo sapiens 26S proteasome-associated ped1 homolog (POH1) mRNA.	Human transposon-like element, partial	Human calcium-dependent phospholipid-binding protein (PLA2) mRNA, complete cds	Human calcium-dependent phospholipid-binding protein (PLA2) mRNA, complete cds	zv90c05.r1 Soares_NhHMPu_S1 Homo sapiens cDNA clone IWAGE:767048 5	RC4-BT0311-141199-011-h06 BT0311 Homo sapiens cDNA	wn57h07.x1 NCI_CGAP_Lu19 Homo sapiensicDNA clone IMAGE:2449597 3' similar to WP:F15G9.4A	CE18595;	wn57h07.x1 NCI_CGAP_Lu19 Homo sapiens cDNA done IMAGE:2449597 3' similær to WP:F15G9.4A CE18595 ;	wn57h07.x1 NCI CGAP Lu19 Homo sapiens cDNA clone IMAGE:2449597 3' similar to WP:F15G9.4A	CE18595;	H.sapiens mRNA for ribosomal protein L31	RC4-BT0311-141199-011-h06 BT0311 Homo saplens cDNA	Homo saplens thyroid hormone receptor binding protein (AIB3), mRNA	Mus musculus fregile X mental retardation syndrome 1 homolog (Fmr1), mRNA	H.saplens DNA for endogenous retroviral like element	Homo sapiens germ-line DNA upstream of Jkappa locus	Human endogenous retrovirus, complete genanie	Homo sapiens methylene tetrahydrofolate dehydrogenase (NAD+ dependent), methenyltetrahydrofolate	cyclohydralase (MTHFD2), mRNA	QV1-DT0069-110200-067-910 DT0069 Homo sapiens cDNA	EST377546 MAGE resequences, MAGI Homo sapiens cDNA	Homo sapiens cAMP-regulated guanine nucleotide exchange factor I (cAMP-GEFI) mRNA, complete cds	
15 P	lop Hit Database Source	T_HUMAN				NT	LN L			IN		TN TN	П	EST_HUMAN I		EST_HUMAN	EST HUMAN	Т	T_HUMAN		T_HUMAN			IN						EST_HUMAN	INT TN	
	Top Hit Acession No.	AI621017.1	11418041 NT	11418322 NT	11418248 NT	AL160311.1	AL160311.1	5031980 NT	5031980 NT	M87299.1	M72393.1	M72393.1		BE064410.1		AI924653.1	A1924653.1		AI924653.1	X69181.1	BE064410.1	11420557 NT	6679816 NT	X89211.1	AJ223364.1	9635487 NT		8843		AW965473.1	U78168.1	
Most Similar	(Top) Hit BLAST E Value		1.0E-65	1.0E-85	1.0E-65	9.0E-68	9.0E-68 ⊅	9.0E-68	9.0E-68	9.0E-66	9.0E-69 N		_	7.0E-68		6.0E-68	6.0E-86	_	6.0E-66		5.0E-66	5.0E-66	4.0E-68	4.0E-66		4.0E-66				4.0E-66	4.0E-66 L	_
	Expression Signal	2.35	2.27	4.85	1.44	1.51	1.51	2.49	2.49	4.18	0.9	6.0	0.88	1.73		1.22	1.22		1.22	70.7	2.25	12.31	0.79	1.94	3.66	5.15		3.35	1.78	4.71	6.89	
	ORF SEQ ID NO:	28681		25318			19865		21097		23529	23530	24268			23944	23945		23946	28636	21107	27424	20542	ZZ0ZZ						24869	26246	l
	SEQ ID NO:	18414	18905	18963	19225	10051	10051		11239	11373	13738	13738	14479	18543		14168	14168		14168	18372	11250	17225	10703	12120	12302	14560		1		15108	16098	ł
1	SEQ ID	8542	9165	9264	9865	65	65	1332	1332	1468	3826	3826	4591	8654		4269	4269		4269	8499	1344	7357	773	2235	2425	4674		5407	5512	6098	6232	

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Probe SEQ ID NO:	SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
8052	17943	28193	1.96	4.0E-66	BF507493.1	EST_HUMAN	UI-H-BW1-amr-a-10-0-UI.s1 NCI_CGAP_Sub7 Homo sapiens cDNA clone IMAGE:3070747 3'
1408	11313	21175	24.62	3.0E-66	4502098 NT	TN	Homo sapiens solute carrier family 25 (mitochondrial carrier; adenins nucleotide translocator), member 5 (SLC25A5), nuclear gene encoding mitochondrial protein, mRNA
1408	11313	21176	24.62	3.0E-66	450209B	١	Homo sapiens solute carrier family 25 (mitochondrial carrier, adenins nucleotide translocator), member 5 (SLC25A5), nuclear gene encoding mitochondrial protein, mRNA
1939	11834	21717	0.84	3.0E-66	N55323.1	EST_HUMAN	yzztg12.r1 Soares, multiple, sclerosis, ZNbHMSP Homo saplens cDNA clane IMAGE:284326 5' similar to SW:H2B1_TIGCA P35068 HISTONE H2B.1/H2B.2. [2] PIR:B56612;
1939	11834	21718	0.84	3.0E-66	N55323.1	EST_HUMAN	yz7g12.r1 Soares_multiple_sclerosis_2NbHMSP Homo saplens cDNA clone IMAGE:284326 5' similar to SW:H2B1_TIGCA P35068 HISTONE H2B.1/H2B.2. [2] PIR:B56612;
1939	11834	21719	0.84	3.0E-66	N55323.1	EST_HUMAN	yz27g12.r1 Şoares_multiple_solerosis_2NbHMSP Homo saplens cDNA clone IMAGE:284326 5' similar to SW:H2B1_TIGCA P35068 HISTONE H2B.1/H2B.2. [2] PIR:B56612;
2675	12540	22430	2.91	3.0E-66	11141880 NT	FZ	Homo sapiens TGF(beta)-induced transcription factor 2 (TGIF2), mRNA
3079	13008	22797	5.47	3.0E-66	7662223 NT	NT	Homo sapiens KIAA0649 gene product (KIAA0649), mRNA
5528	15445	25511	1.64	3.0E-66	11417946 NT	NT	Homo sapiens NIPSNAP, C. elegans, homolog 1 (NIPSNAP1), mRNA
6528	16445	25512	1.84	3.0E-66	11417948 NT	NT	Homo sapiens NIPSNAP, C. elegans, homolog 1 (NIPSNAP1), mRNA
8800	18614	28904	8.3	3.0E-66	5453949 NT	۲	Homo sapiens protein phosphatase 2, regulatory subunit B (B56), alphe isoform (PPP2R5A) mRNA
45	10033	19837	1.02	2.0E-66	7657334 NT	TN	Homo sapiens Misshapen/NiK-related kinase (MiNK), mRNA
45	10033	19838	1.02	2.0E-66	T657334 NT	TN	Homo saplens Misshapen/NIK-related kinase (MINK), mRNA
416	8883	19774	6.0	2.0E-66	4505524 NT	١	Homo saplens origin recognition complex, subunit 5 (yeast homolog)-like (ORCSL) mRNA, and translated products
418	9983	19775	0.93	2 0F-88	4505524 NT	L	Homo seplens origin recognition complex, subunit 5 (yeast homolog)-like (ORC5L) mRNA, and translated products
1784	L			2.0E-68	AL163301.2	NT	Homo sapiens chromosome 21 segment HS21C101
2944	12871	22669	0.95	2.0E-66	X65859.1	N	H.sapiens pseudogene for the low affinity IL-8 receptor
3975	13882	23657	0.88	2.0E-86	AF108389.1	NT	Homo sapiens sodium/calcium exchanger isoform NaCa3 (NCX1) mRNA, complete cds
4656	14448	24233	12.69	2.0E-66	AJ133267.2	NT	Homo saplens HLA-B gene for human leucocyte antigen B
4556	li			2.0E-66	AJ133267.2	NT	Homo sapiens HLA-B gene for human leucocyte antigen B
7135	17012	27205	2.16	2.0E-66	N45480.1	T HUMAN	yy59c02.r1 Soares_multiple_sclerosis_2NbHM3P Homo sapiens cDNA clone IMAGE:277826 5'
9475				2.0E-66	11418318 NT	NT	Homo sapiens G-2 and S-phase expressed 1 (GTSE1), mRNA
2864	12792		1.38	1.0E-66	AV717817.1	EST_HUMAN	AV717817 DCB Hamo sapiens cDNA clone DCBADC07 5'
2864	1	22586					AV717817 DCB Homo sapiens cDNA clone DCBADC07 5'
4288	- 1					EST_HUMAN	AV717817 DCB Hamo sapiens cDNA clone DCBADC07 5'
4288	12792	22586	3.26	1.0E-66	AV717817.1	EST_HUMAN	AV717817 DCB Hamo sapiens cDNA clone DCBADC07 6'

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Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession . No.	Top Hit Database Source	Top Hit Descriptor
5306		25031	5.36	1.0E-66		EST_HUMAN	602152996F1 NIH_MGC_81 Homo sapiens cDNA clone IMAGE:4294161 5'
6143	15991	26126	1.49	1.0E-66		EST_HUMAN	RC5-BN0193-010800-034-G06 BN0193 Homo sapiens cDNA
6926	16804	26998	1.37	1.0E-68	1.0E-66 AA668858.1	EST_HUMAN	aa80e04.s1 NCI_CGAP_GCB1 Homo saplens cDNA clone IMAGE:827262 3'
8312		28438	2.39	1.0E-68		LN LN	Homo sapiens jun dimerization protein gene, partial cds; cfos gene, complete cds; and unknown gene
9260	18985		2.09	9.0E-67	9.0E-67 11418177 NT	TN	Homo saplens Ran GTPase activating protein 1 (RANGAP1), mRNA
4829	•		28.0	8.0E-67	8.0E-67 M78158.1	EST_HUMAN	EST01750 Subtracted Hippocampus, Stratagene (cat. #936205) Homo sapiens cDNA clone HHCPN31 similar to L1 repetitive element
376	10360	20184	6.19	7.0E-67	7.0E-67 AW 162232.1	EST_HUMAN	au75d02.x1 Schneider fetal brain 00004 Homo sapiens cDNA clone IMAGE:2782083 3' similar to gb:M37104 ATP SYNTHASE COUPLING FACTOR 6, MİTOCHONDRIAL PRECURSOR (HUMAN);
1360	11288	21122	2.63	7.0E-87	7.0E-67 AA383416.1	EST_HUMAN	EST98812 Testis I Homo sapiens cDNA 5' erid similar to similar to C. elegans hypothetical protein, cosmid ZK353
1535	ı	21298	-	7.0E-67		EST_HUMAN	zh56b05.r1 Soares_fetal_liver_spleen_1NFLS_S1 Homo saplens cDNA clone IMAGE:416049 5'
1536	11439	21297	+	7.0E-67	7.0E-67 W85947.1	EST_HUMAN	zh56b05.r1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA done IWAGE:416049 5'
1988	11881	21773	1.06	7.0E-67	7657243 NT	NT	Homo sapiens inositol 1,3,4-triphosphate 5/8 kinase (ITPK1), mRNA
1988	11881	21774	1.08	7.0E-67	7657243 NT	NT	Homo saplens Inositol 1,3,4-triphosphate 5/6 kinase (ITPK1), mRNA
2779		20184	7.07	7.0E-67	7.0E-67 AW 162232.1	EST_HUMAN	au76d02.x1 Schneider fetal brain 00004 Home saplens cDNA clone IMAGE:2782083 3' similar to gb:M37104 ATP SYNTHASE COUPLING FACTOR 6, MITOCHONDRIAL PRECURSOR (HUMAN);
5793		25807	2.04	7.0E-67	11425572 NT	LΝ	Homo sapiens adaptor-related protein complex 2, beta 1 subunit (AP2B1), mRNA
6793		25808	2.04	7.0E-67	11425572 NT	NT	Homo sapiens adaptor-related protein complex 2, beta 1 subunit (AP2B1), mRNA
8038		29111	1.56	7.0E-67		NT	Homo sapiens low density lipoprotein-related protein 2 (LRP2), mRNA
8038			1.56	7.0E-67	11430460 NT	LN	Homo sapiens low density lipoprotein-related protein 2 (LRP2), mRNA
9501		25292	3.33	7.0E-67 A	AB0113	NT	Homo sapiens gene for AF-6, complete cds
9868	19357		1.43	7.0E-67	11421527 NT	NT	Homo sapiens calcium channel, voltage-dependent, alpha 2/delta subunit 1 (CACNA2D1), mRNA
547			1.32	6.0E-67	6.0E-67 X68968.1	NT	H.saplens mRNA for acetyl-CoA carboxylase
778	10708	20247	1.5		6.0E-67 Z17227.1	IN	Homo sapiens mRNA for transmebrane receptor protein
1252			0.93	6.0E-67	Y14320.1	LN	Homo sapiens PMP69 gene, exons 3,4,5,6 & 7
3131	13058		1.24	6.0E-67	4506434 NT	NT	Homo saplens retinoblastoma 1 (including osteosarcoma) (RB1) mRNA
3391	13308		1.2	6.0E-67	4507332 NT	NT	Homo saplens Synapsin III (SYN3) mRNA, and translated products
3391	13308	23107	1.2	6.0E-67	4507332 NT	NT	Homo sapiens Synapsin III (SYN3) mRNA, and translated products
4035	13938		1.28	6.0E-67 AI	.163201.2	NT	Homo saplens chromosome 21 segment HS210001
4035		23715		6.0E-67 A	L163201.2	NT	Homo sapiens chromosome 21 segment HS210001
4607			3.37	6.0E-67		NT	Homo sapiens DKFZp434P211 protein (DKFZP434P211), mRNA
4607	14495	24284	3.37	6.0E-67	7657020 NT	NT	Homo saplens DKFZp434P211 protein (DKFZP434P211), mRNA

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
5125	14761		2.1	6.0E-67	4507848 NT	TN	Homo sapiens ubiquitin specific protease 13 (isopeptidase T-3) (USP13) mRNA
3184	13109	22913	2.45	5.0E-67	5.0E-67 AF009660.1	NT	Homo sapiens T cell receptor beta locus, TCRBV7S3A2 to TCRBV12S2 region
8362	18229		2.1	5.0E-67	5.0E-67 BE010038.1	EST_HUMAN	PM3-BN0176-100400-001-g04 BN0176 Homo sapiens cDNA
1306	11213	21069			4.0E-67 R90819.1	EST_HUMAN	yn02d11,r1 Soares adult brain N2b4HB55Y Homo saplens cDNA clone IMAGE:167253 5
6883	16762		1.22		4.0E-67 BF357321.1	EST_HUMAN	RC0-HT0934-150900-026-c03 HT0934 Homo saplens cDNA
8416	. 18290		2.3	4.0E-67	AA714294.1	EST_HUMAN	nw06e01.s1 NCI_CGAP_SS1 Homo saplens cDNA clone IMAGE:1238472 3' similar to TR:O10385 O10385 PRO-POL-DUTPASE POLYPROTEIN;
2782	10553	20365	0.93	3.0E-67	3.0E-67 AA333768.1	EST_HUMAN	EST37903 Embryo, 9 week Homo sapiens cDNA 5' end
3407	13324	23125	1.14	3.0E-67	BE064410.1	EST_HUMAN	RC4-BT0311-141199-011-h06 BT0311 Homo sapiens cDNA
4596	14484	24270	3.14	3.0E-67	3.0E-67 AW869159.1	EST_HUMAN	MR3-SN0066-040500-008-f01 SN0066 Homo sapiens cDNA
6760	16639	26827			3.0E-67 BF196068.1	EST_HUMAN	hr81f05.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:3134913 3' similar to SW:RHOP_MOUSE Q61085 GTP-RHO BINDING PROTEIN 1;
8583	18451		19.27		3.0E-67 AA927874.1	EST_HUMAN	om18b07.s1 Soares_NFL_T_GBC_S1 Homo sepiens cDNA clone IMAGE:1541365 3'
180	10152	18967			2.0E-67 BE348354.1	EST HUMAN	hw16g09.x1 NCI_CGAP_Lu24 Homo sepiens cDNA clone IMAGE:3183136 3' similar to WP:F23H11.9 CE09617
827	10754			2.0E-67	AW816405.1	EST HUMAN	QV4-ST0234-181199-037-f05 ST0234 Homo(saplens cDNA
1089	11005		1.74	2.0E-67	2.0E-67 AF167460.1	닏	Homo sapiens double stranded RNA activated protein kinase (PKR) gene, exons 2a, 2, 3, and 4
1841	11737	21614	1.5		2.0E-67 BE303037.1	EST_HUMAN	ba72g05.y1 NIH_MGC_20 Homo sapiens cDNA clone IMAGE:2905976 5' similar to TR:094892 094892 KIAA0798 PROTEIN .
1841	11737	21615	1.5		2.0E-67 BE303037.1	EST_HUMAN	ba72g05.y1 NIH_MGC_20 Homo sapiens cDNA clone IMAGE:2905976 5' simitar to TR:O94892 O94892 KIAA0798 PROTEIN ;
2336	12216	22116	86.0	2.0E-67 AF	AF309561.1	F	Homo saplens KRAB zinc finger protein ZFQR mRNA, complete cds
2381	12281	22153	1.2	2.0E-67	4758795 NT		Homo saplens developmentally regulated GTP binding protein 1 (DRG1), mRNA
3422	13339	23144	3.9		2.0E-67 AA625755.1	EST_HUMAN	zu91g01.s1 Soares_testis_NHT Homo sapient cDNA clone IMAGE:7453923'
3921	13830			2.0E-67 AL	AL163300.2	NT	Homo sapiens chromosome 21 segment HS21C100
5724	15631	75734	4.22	2.0E-67	2.0E-67 BF240758.1	EST_HUMAN	601875351F1 NIH_MGC_55 Homo sapiens cDNA clone IMAGE:4091893 5'
5803	15708	25820	2.17	2.0E-67	2.0E-67 AB051763.1	NT	Homo sapiens mRNA for NADPH-cytochrome P-450 reductase, complete cds
5803	15708	25821	2.17	2.0E-67	2.0E-67 AB051763.1	NT	Homo saplens mRNA for NADPH-cytochrome P-450 reductase, complete cds
7202				2.0E-67	2.0E-67 AW602635.1	EST_HUMAN	RC4-BT0566-170100-011-c07 BT0566 Homo saplens cDNA
7202		27265	1.34	2.0E-67	2.0E-67 AW602635.1	EST_HUMAN	RC4-BT0566-170100-011-c07 BT0566 Homo sapiens cDNA
8409	19769		3.26	2.0E-67	11436448 NT	TN	Homo sapiens KIAA0985 protein (KIAA0985), inRNA
8558					BE295714.1	EST_HUMAN	601175762F1 NIH_MGC_17 Homo sapiens cDNA done IMAGE:3531038 5'
8751		28144	,,		2.0E-67 BF377169.1	EST_HUMAN	PM2-TN0103-040900-001-c02 TN0103 Homo sapiens cDNA
9388	19577	╛	2.6		11418189 NT	뉟	Homo sepiens thyroid autoantigen 70kD (Ku antigen) (G22P1), mRNA
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gt38h02.x1 Soares_fetal_lung_NbHL19W Homo sapiens cDNA clone IMAGE:1950291 3' similar to contains Homo sapiens killer inhibitory receptor 2-2-1 (KIR221) and killer inhibitory receptor 2-2-2 (KIR222) genes, Homo sapiens amyloid beta (A4) precursor protein (protease nexin-II, Alzheimer disease) (APP), mRNA zq82h10.r1 Stratagene hNT neuron (#837233) Homo sapiens cDNA clone IMAGE:648163 5' similar to zq8Zh10.r1 Stratagene hNT neuron (#937233) Homo sapiens cDNA clone IMAGE:648163 5' similar to UI-HF-BN0-elb-c-07-0-UI.r1 NIH_MGC_60 Homo sepiens cDNA clone IMAGE:3078924 5 Homo sepiens brefeldin A-inhibited guanine riucleoilde-exchange protein 2 (BIG2), mRNA yg38g04.s1 Soares Infant brain 1NIB Homo sepiens cDNA clone IMAGE:34896 3' vlus musculus G-protein coupled receptor GPR73 (Gpr73) mRNA, complete cds 601452067F1 NIH_MGC_66 Homo sapiens cDNA clone IMAGE:3855761 5 601894635F2 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:4124144 5 Homo sepiens serine carboxypeptidase 1 precursor protein (HSCP1), mRNA Homo sapiens serine carboxypeptidase 1 precursor protein (HSCP1), mRNA 65 Homo sapiens cDNA clone IMAGE:3852254 5 GLYCERALDEHYDE 3-PHOSPHATE DEHYDROGENASE, LIVER **Fop Hit Descriptor** Homo sapiens gene for activin receptor type IIB, complete cds Cricetulus longicaudatus mRNA for EF-1 alpha, complete cds QV1-DT0072-010200-056-h08 DT0072 Homo sapiens cDNA HSPD18178 HM3 Homo sapiens cDNA clone s3000023D09 Homo sapiens mRNA for KIAA0145 protein, partial cds Homo saplens mRNA for KIAA0145 protein, partial cds Homo sapiens mRNA for KIAA1485 protein, partial cds Homo sapiens mRNA for KIAA1431 protein, partial cds Homo sepiens transcription factor NRF (NRF), mRNA Homo sepiens transcription factor NRF (NRF), mRN/ Homo sapiens chromosome 21 unknown mRNA Homo sapiens chromosome 21 unknown mRNA Homo sapiens chromosome 21 unknown mRNA Homo sapiens chromosome 21 unknown mRINA SW:SAV_SULAC Q07590 SAV PROTEIN. SW:SAV_SULAC Q07590 SAV PROTEIN. THR.12 THR repetitive element; partial ods EST_HUMAN EST_HUMAN **EST HUMAN** EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN **EST HUMAN** EST_HUMAN HUMAN Top Hit Database SWISSPROT Source EST A 눋 눌 Ę 뉟뉟 与 4502166 NT 눋 11422086|NT 11421388 11421388 Top Hit Acession 11055991 4.0E-68 D63479.2 4.0E-68 AB040918.1 3.0E-68 AF236082.1 3.0E-68 AW939485.1 6.0E-68 AW 503842.1 2.0E-68 AB008681.1 8.0E-68 AA209456.1 6.0E-68 AF133801.1 6.0E-68 BE612554.1 6.0E-68 BF310676.1 5.0E-68 AF231919.1 5.0E-68 AF231919.1 3.0E-68 AI342323.1 BE870732.1 AA209456.1 5.0E-68 AF231919.1 5.0E-68 AF231919.1 5.0E-68 AB037852.1 ġ 3.0E-68 F28784.1 2.0E-68 D00522.1 4.0E-68 4.0E-68 2.0E-68 4.0E-68 4.0E-68 1.0E-67 8.0E-68 8.0E-68 4.0E-88 6.0E-88 (Top) Hit BLAST E Aost Simila Value 0.87 0.87 3.87 5.64 1.53 1.66 8 2 4.44 4.37 2.46 2.48 8 1.42 3.87 2.62 1.01 5. 5.64 5.41 5.61 4.96 Expression Signal 20036 20556 20572 22230 22248 26189 28088 23492 23483 20573 27290 24261 28627 20555 26188 28064 27291 ORF SEQ ÖNO 17847 19502 15076 14473 10219 16454 18362 12356 12356 16044 16044 13525 13706 13706 12642 SEQ ID 12017 17821 19234 10731 10731 11745 1939 Š ÿ 7897 9872 2832 4583 6110 7974 3108 4900 2480 7441 253 2129 3784 3794 8489 9876 9918 785 802 298 6061 225 SEO ID 28 785 2480 6061 7225 3611 ÿ

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Top Hit Descriptor	601458514F1 NIH_MGC_66 Homo sapiens cDNA clone IMAGE:3862034 5	601437367F1 NIH_MGC_72 Homo sepiens cDNA clone IMAGE:3922192 5	Homo sapiens meningioma (disrupted in balanced translocation) 1 (MN1), mRNA	QV4-ST0234-181199-037-f05 ST0234 Home sapiens cDNA	Homo sapiens mRNA for KIAA0577 protein, complete cds	Homo sapiens mRNA for KIAA0577 protein, complete cds	ULH-Bi3-alk-f-01-0-UI.s1 NCI_CGAP_Sub5 Homo sapiens cDNA clone IMAGE:273727233	601177002F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:3532344 5'	al47g12.s1 Soares_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:1460518 3'	Homo sapiens cell recognition molecule Caspi2 (KIAA0868), mRNA	Homo sapiens phosphodiesterase 7B (PDE7E), mRNA	Homo sapiens phosphodiesterase 7B (PDE7B), mRNA	Homo sapiens MIF2 suppressor (HSMT3) mRNA, complete cds	Homo sapiens meningioma (disrupted in balanced translocation) 1 (MN1), mRNA	Homo sapiens low density lipoprotein-related protein 2 (LRP2), mRNA	Homo sapiens pre-B-cell colony-enhancing factor (PBEF) mRNA	Homo sapiens pre-B-cell colony-enhancing factor (PBEF) mRNA	Homo sapiens 26S proteasome-associated pad1 homotog (POH1) mRNA	Homo sapiens 26S proteasome associated pad1 homolog (POH1) mRNA	Homo saplens v-raf murine sarcoma viral oncosene homolog B1 (BRAF) mRNA	AU117241 HEMBA1 Homo sapiens cDNA clore HEMBA1000968 5'	Homo sapiens RIBIIR gene (partial), exon 12	Homo sapiens actin-related protein 3-beta (ARP3BETA), mRNA	qe82h01.x1 Soares_fetal_lung_NbHL19W Horio saplens cDNA clone IMAGE:1743601.3' sImilar to gb:L11566 60S RIBOSOMAL PROTEIN L18 (ḤUMAN);	qe62h01.x1 Soares_fetal_lung_NbHL19W Homo sapiens cDNA clone IMAGE:1743601.3' similar to	wm28h11x1 NCI CGAP Ut4 Homo sepiens cDNA clone IMAGE:2437125 3*	wh57b08.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2384819 3' similar to TR:055137	O66137 ACYL-COA THIOESTERASE.;	Homo sapiens latent transforming growth factor, beta binding protein 2 (LTBP2) mRNA	Homo sapiens latent transforming growth factor beta binding protein 2 (LTBP2) mRNA	601110371F1 NIH_MGC_16 Homo capiens cDNA clone IMAGE:3361352 6'	Homo sapiens Smad- and Olf-Interacting zinc finger protein mRNA, partial cds
Top Hit Database Source	EST_HUMAN	EST_HUMAN	IN	EST_HUMAN	IN	TN	EST_HUMAN	EST_HUMAN	EST_HUMAN	IN	N-	٦	Ŋ	N	NT TN	NT	NT	NT	IN	NT	EST_HUMAN	IN	IN	EST_HUMAN	MAN 12 HOR	EST HUMAN		EST_HUMAN	TN	NT	EST_HUMAN	NT
Top Hit Acession No.	BF035316.1	BE897376.1	4505222 NT	1.0E-68 AW816405.1	1.0E-68 AB011149.1	1.0E-68 AB011149.1	1.0E-68 AW451832.1	1.0E-68 BE296032.1	1.0E-68 AA897343.1	7682349 NT	11418869 NT	11418869 NT	L76416.1	4505222 NT	11430460 NT	5031976 NT	5031976 NT	5031980 NT	5031980 NT	4757867 NT	AU117241.1	8.0E-69 AJ237744.1	JN 2169966.	6.0E-69 AI192764.1	A1400764 4	4.0E-69 AIR73630.1		4.0E-69 AI764973.1	4557732 NT	4557732 NT		3.0E-69 AF221712.1
Most Similar (Top) Hit BLAST E Value	2.0E-68 BF	2.0E-68	1.0E-68	1.0E-68				1.0E-68	1.0E-68	1.0E-88	1.0E-68	1.0E-68	1.0E-68	1.0E-68	1.0E-68	9.0E-69	9.0E-69	9.0E-69	9.0E-69	9.0E-69	9.0E-69	8.0E-69	7.0E-69	6.0E-69	900	4 0F-69		4.0E-69	4.0E-69	4.0E-69	3.0E-69 B	3.0E-69
Expression Signal	4.79	.1.92	1.31	9.36	1.32	1.32	1.01	0.95	96.0	1.57	2.44	2.44	2.29	1.95	1.38	1.82	1.82	1.63	1.63	0.78	9.27	1.21	80'9	3.61	6	188		4.12	2.43	2.43	2.81	1.64
ORF SEQ ID NO:	26095		19873	20078	21894			23617	24598	24929	28361	28362	28396	19873	24991	19797	19798		20773	23718			25852	26659	26860			25554	26024	26025		20340
Exan SEQ ID NO:			10056	10257	12092	12092		13837	14831		18108	18108	18155	10056		10006	10008		10929	13940	18143	13260	15740	16469		10451		16481	15900			10532
Probe SEQ ID NO:	6202	9148	72	293	2205	2205	2728	3928	4954	5261	8226	8228	8275	6996	9862	19	19	1011	1011	4037	8283	3340	5834	6588	6580	208		5565	5895	2882	380	969

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Top Hit Descriptor	yd08a02.r1 Soares infant brain 1NIB Homo sepiens cDNA clone IMAGE:24880 5' similar to SP:A48836 A48836 SPEGF III=EGF REPEAT-CONTAINING FIBROPELLIN-LIKE PROTEIN - SEA URCHIN ;	Homo sapiens lymphatic vessel endothelial hyaluronan receptor 1 (LYVE-1) mRNA	wh68g08.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2385768 3'	Homo sapiens aconitase 2, mitochondrial (ACO2), mRNA	Homo sapiens arm-repeat protein NPRAP/neurojungin (CTNND2) mRNA, partial de	Homo sapiens TRAF6-binding protein T6BP mRNA, complete cds	EST88807 HSC172 cells II Homo saplens cDNA 5' end similar to similar to ribosomal protein S18	H.sapiens mRNA for N-acetylglucosamide (beta 1-4)-galactosyltransferase	Human mRNA for calcium-binding protein in macrophages (MRP-14) macrophage migration inhibitory factor	(MIF)-related protein	Homo sapiens ribosomal protein S15a (RPS15A), mRNA	EST88807 HSC172 cells II Homo sapiens cDNA 5' end similar to similar to ribosomal protein S18	Homo sapiens HGC6.2 protein (HGC6.2), mRNA	Homo sapiens KIAA0553 protein gene, complete cds, and alphalib protein gene, partial cds	Homo sapiens KIAA0553 protein gene, complete cds; and alphallb protein gene, partial cds	Homo sapiens KIAA0553 protein gene, complete cds, and alphallb protein gene, partial cds	Homo sapiens KIAA0553 protein gene, complete cds; and alphallb protein gene, partial cds	601109444F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3350074 5'	zw71g02.r1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:781682 5	Rattus norvegicus brain specific cortactin-binding protein CBP90 mRNA, partial cds	QV0-TT0010-031199-045-c07 TT0010 Homo sapiens cDNA	Homo sapiens KIAA0716 gene product (KIAA0716), mRNA	Homo sapiens KIAA0716 gene product (KIAA0716), mRNA	Homo sapiens mRNA for KIAA1147 protein, partial cds	Homo saplens mRNA for KIAA1147 protein, partial cds	TCBAP1E2678 Pediatric pre-B cell acute lymphoblastic leukemia Baylor-HGSC project=TCBA Homo sapiens	cDNA clone TCBAP2678	TCBAP1E2678 Pediatric pre-B cell acute lymphoblastic leukemia Baylor-HGSC projecte TCBA Homo sapiens	CLINA date 1 CDAF 2076	Homo sapiens Keraiin 8 (KK18) mKNA	601762902F1 NIH_MGC_20 Homo saplens cDNA clone IMAGE:4025785 5
Top Hit Database Source	EST_HUMAN	FZ	EST_HUMAN	NT	NT	ΙN	EST_HUMAN	TN	!	LN	ΝT	EST_HUMAN	NT	NT	NT	IN	NT	EST_HUMAN	EST_HUMAN	NT	EST_HUMAN	NT	NT	LN T	١		EST_HUMAN		EST TOWAIN		EST_HUMAN
Top Hit Acession No.	T80514.1		AI765888.1	11418185 NT	J52351.1		9.1				11432120 NT	3.0E-69 AA376399.1	11419157 NT	2.0E-69 AF160252.1	2.0E-69 AF160252.1		2.0E-69 AF160252.1	2.0E-69 BE257857.1	2.0E-69 AA431157.1	1.0E-69 AF053768.1	1.0E-69 AW393969.1	7662263 NT	7662263 NT	1.0E-69 AB032973.1	1.0E-69 AB032973.1		1.0E-69 BE245070.1		1.0E-09 BEZ45070.1	4504918 N	3F125887.1
Most Similar (Top) Hit BLAST E Value	3.0E-69	3.0E-69	3.0E-69 A	3.0E-69	3.0E-69 U	3.0E-69	3.0E-69	3.0E-69 X13223.1		3.0E-69 X06233.1	3.0E-69	3.0E-69	3.0E-69	2.0E-69	2.0E-69/	2.0E-69	2.0E-69 /	2.0E-69	2.0E-69	1.0E-69	1.0E-69	1.0E-69	1.0E-69	1.0E-69	1.0E-69/		1.0E-69		1.05-09 (1.0E-69	1.0E-69 B
Expression Signal	1.35	0.88	,0.86	5.94	1.37	8.43	1.28	1.54		2.24	3.07	7.12	4.13	1	 	4.94	4.94	1.2	2.73	2.35	3.68	1.55	1.55	2.93	2.83		5.29		87.0	23.2/	1.63
ORF SEQ ID NO:			23874	29105				17491			28174			20170	20171		20171	21616		21445	26008	26211		26173	26174		27942		2/943		28785
Exon SEQ ID . NO:	11440				16249	16318	16968	17284		- 1		18099			10344	10344	10344				15886	16063		16033	16033		17697		_L	┙	18872
Probe SEQ ID NO:	1536	2325	3990	5209	6387	6457	7091	7417		7483	8038	8215	9168	124	124	388	398	1842	2813	1675	5981	0809	6080	8809	6088		7847		/86/	8240	9105

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Single Exon Probes Expressed in Heart

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	Top Hit Descriptor	wf64e08.x1 Scares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2360390 3' similar to contains Alurepetitive element; contains element MIR repetitive element;	nc13d12.r1 NCI_CGAP_Pr1 Homo sapiens cDNA clone IMAGE:1008023	Homo sapiens DGS-I mRNA, 3' end	tm89f01.x1 NCI_CGAP_Bm25 Homo sapiens cDNA clone IMAGE:2165305 3'	tm89f01.x1 NCI_CGAP_Brn25 Hano sapiens cDNA clone IMAGE:2165305 3'	Z15h04.r1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:713239 5'	Homo sapiens tumor suppressor deleted in oral cancer-related 1 (DOC-1R) mRNA	Homo sapiens adenylate cyclase 3 (ADCY3) mRNA	Homo sapiens MIST mRNA, partial cds	Homo sapiens MIST mRNA, partial cds	Homo sapiens gene encoding splicing factor SF1, exons 2-8	Homo sapiens mRNA for KIAA1294 protein, partial cds	Homo sapiens mRNA for KIAA1294 protein, partial cds	Human displacement protein (CCAAT) mRNA	Human displacement protein (CCAAT) mRNA	Human PBX3 mRNA	Human PBX3 mRNA	Horno sapiens phospholipid scramblase 1 gene, exon 1 and 5' flanking region	Homo sapiens karyopherin beta 2b, transportin (TRN2), mRNA	Homo sapiens karyopherin beta 2b, transportin (TRN2), mRNA	Homo sapiens HIR (histone cell cycle regulation defective, S. cerevisiae) homolog A (HIRA), mRNA	Homo sapiens HIR (histone cell cycle regulation defective, S. cerevisiae) homolog A (HIRA), mRNA	Homo saplens amyloid beta (A4) precursor protein (protease nexin-II, Alzheimer disease) (APP), mRNA	Human Ku (p70/p80) subunit mRNA, complete cds	Homo saplens CMP-N-acetylneuraminic acid synthase (LOC55907), mRNA	Homo saplens KJAA0792 gene product (KIAA0792), mRNA	Homo sapiens KJAA0792 gene product (KIAA0792), mRNA	MR3-HT0487-150200-115-a06 HT0487 Homo saplens cDNA	RC0-BT0522-071299-011-a12 BT0522 Homo sapiens cDNA	RC0-BT0522-071299-011-a12 BT0522 Homo sapiens cDNA
Top Hit	Database	EST_HUMAN	EST_HUMAN	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	IN	IN	TN	۲	NT	NT	NT	NT	IN	۲N	NT	TN	IN	IN	TN	LN	ΓN	NT	IN	IN	IN	EST_HUMAN	EST_HUMAN	EST_HUMAN
Top Hit Acession	No.	4/809994.1	4A230303.1	_77566.1	41497807.1	41497807.1	4A282955.1	5031668 NT	4757723 NT	4B032369.1	4B032369.1	4J000052.1	AB037715.1	4B037715.1	M74099.1	M74099.1	X59841.1	X59841.1	AF153715.1	11525964 NT	11525964 NT	11526319 NT	11526319 NT	4502166 NT	M30938.1	8923899 NT	7862307 NT	7662307 NT	3E166034.1	3E071796.1	3E071786.1
Most Similar	BLAST E	1.0E-69	8.0E-70	8.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	8.0E-70	6.0E-70	6.0E-70	5.0E-70	5.0E-70	5.0E-70	3.0E-70 B	3.0E-70 B
Everescion	Signal	2.32	1.52	2.16	1.93	1.93	1.83	3.57	3.88	5.28	5.28	1.89	2.36	2.36	3.98	3.98	3.72	3.72	3.13	1.56	1.56	1.78	1.78	1.83	1.36	0.39	1.83	1.83	1.73	0.89	0.89
CHA	Ö ÖÖ:		22065	23954	21547	21548	21660		23807	25124	25125		26980	26981	27131		27357	27358	26627	26848	26649	28978	28979	20630			22268	52269	.	21332	
Exon	850 80 .00	19124	12717	14176	11670	11670			14032	15289		15985	16788	16788	16940		17159			16457	16457	18687	18687	10780	11979		12723	12723		11475	11475
Probe	SEO ID	9510	2284	4277	1771	1771	1888	2018	4132	5369	6369	6138	6910	6910	5907	7063	7283	7283	7428	7445	7445	8875	8875	853	2090	2481	2505	2505	9116	1571	1571

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Top Hit Descriptor	602141561F1 NIH_MGC_46 Homo sapiens :DNA clone IMAGE:4302806 5'	602141561F1 NIH_MGC_46 Homo sapiens cDNA clone IMAGE:4302806 5'	ую7а10.r1 Soares melanocyte 2NbHM Homo septens cDNA clone IMAGE:270522 5' similar to SW:D3HI_RAT P29268 3-HYDROXYISOBUTYRATE DEHYDRÓGENASE PRECURSOR;	yo7a10.r1 Soares melanocyte 2NbHM Home sapiens cDNA clone IMAGE:270522 5' similar to SW:D3HI RAT P29288 3-HYDROXYISOBUTYRATE DEHYDROGENASE PRECURSOR	qx51h01 x1 NCL CGAP_Pan1 Homo sapiens cDNA clone IMAGE:2004913 3'	Homo saplens hypothetical protein FLJ20758 (FLJ20758), mRNA	Homo sapiens KIAA0193 gene product (KIAA0193), mRNA	Homo sapiens KIAA0193 gene product (KIAA0193), mRNA	hz64c12.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3212758 3'	Homo sapiens chromosome 21 segment HS21C002	z48g04.r1 Soares retina N2b4HR Homo sapiens cDNA clone IMAGE:380214 5' similar to SW:GAG_HTL1A P03345 GAG POLYPROTEIN ;	Human nonmuscle myosin heavy chain-B (MYH10) mRNA, partial cds	H. saplens gene for schwannomin (CS8)	H.sapiens gene for schwannomin (CS8)	Homo sapiens NALP1 mRNA, complete cds	Human mRNA for NF1 protein Isoform (neurofibromin Isoform), complete cds	Homo sapiens cytoplasmic dynein intermediate chain 1 mRNA, complete cds	Homo sapiens cytoplasmic dynein intermediate chain 1 mRNA, complete cds	Homo sapiens stalytransferase 6 (N-acetyllaccsaminide alpha 2,3-sialytransferase) (SIAT6), mRNA	Human guanine nucleotide-binding protein alpha-subunit gene (G-s-aipha), exons 4 and 5	Homo sapiens calcium-binding transporter mRNA, partial cds	Homo sapiens hypothetical protein FLJ20450 (FLJ20450), mRNA	Homo sapiens hypothetical protein FLJ20450 (FLJ20450), mRNA	Homo sapiens eukaryotic translation initiation factor 3, subunit 6 (48kD) (EIF3S6) mRNA	Homo sapiens low density lipoprotein-related protein 2 (LRP2), mRNA	Homo sapiens low density lipoprotein-related protein 2 (LRP2), mRNA	Homo sapiens transglutaminase 3 (E polypeptide, protein-glutamine-gamma-glutamyltransferase) (TGM3) mRNA	zv54c03.r1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:757444 5'	JAV738538 CB Homo sapiens cDNA clone CBLBGB10 5
Top Hit Database Source	EST_HUMAN	EST_HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	NT	Z	NT	EST_HUMAN	ΙN	EST_HUMAN	N	NT .	Ę	NT	M	LN LN	NT	TN	IN	NT	TN	IN	LN	TN	LN		HUMAN	EST HUMAN
Top Hit Acession No.	3.0E-70 BF685233.1	BF685233.1	2.0E-70 N42161.1	N42161.1		TN 6992368	7661983 NT	7661983 NT	2.0E-70 BE467311.1		AA054010.1	2.0E-70 M69181.1	X72662.1	X72662.1	AF310105.1	D12625.1	AF123074.1	2.0E-70 AF123074.1	11422842 NT	2.0E-70 M21741.1	AF123303.1	8923420 NT	8923420 NT	4503520 NT	11430460 NT	11430460 NT	4507476 NT	1.0E-70 AA442292.1	1.0E-70 AV738538.1
Most Similar (Top) Hit BLAST E Value	3.0E-70	3.0E-70	2.0E-70	2.0E-70 N	2.0E-70 A	2.0E-70	2.0E-70	2.0E-70	2.0E-70	2.0E-70	2.0E-70 A	2.0E-70	2.0E-70	2.0E-70	2.0E-70	2.0E-70	2.0E-70	2.0E-70	2.0E-70	2.0E-70	2.0E-70 A	2.0E-70	2.0E-70	2.0E-70	2.0E-70	2.0E-70	1.0E-70	1.0E-70	1.0E-70
Expression Signal	3.88	3.88	13.15	13.15	2.01	1.7	1.95	1.95	0.97	2.09	3.62	3.95	8.05	8.05	1.42	1.88	9.83	9.83	1.69	7.67	1.3	3.19	3.19	5.82	2.58	2.58	2.97	2.57	13.73
ORF SEQ ID NO:	25865	25866	20426	20427		20766	20924	20925	21148	21479		23648	25156	25157	25780	26028	·	26043	24853	26685	27930	28550	28551						28429
Exon SEQ ID NO:	15751	1	10608	10608	L.	10922	11079	11079		11609	12156	L	15304			15904		15915	15134	16498		18296	18296			19114	13267		18182
Probe SEQ ID NO:	5845	5845	674	674	689	1004	1167	1167	1387	1708	2272	3963	5385	5386	5766	6669	6010	6010	6177	6618	7835	8422	8422	8068	9499	9499	3347	7642	8305

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).E		
Probe SEQ ID NO:	SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acessian No.	Top Hit Database Source	Top Hit Descriptor
5621	15536	25621	6.04	9.0E-71	143870.1	EST_HUMAN	qe04f01.x1 Soares_testis_NHT Homo saplens cDNA clone IMAGE:1738009 3' similar to TR:O14045 O14045 PHOSPHOTRANSFERASE: ;
5621	15536	25622	6.04	9.0E-71 AI	143870.1	EST_HUMAN	qe04f01.x1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1738009 3' sImilar to TR:O14045 O14045 PHOSPHOTRANSFERASE.;
6192	16077	26226	1.88		654903.1	EST_HUMAN	wb52c05.x1 NCI_CGAP_GC6 Homo sapiens cDNA clone IMAGE:2309288 3' similar to TR:P97213 P97213 CDU2, CDU1, TCDD, TCDB, TCDE, TCDC, CDD1, CDD2, CDD3, AND CDD4 GENES.
8811	16077	26226	4.65	9.0E-71 AI	A1654903.1	EST_HUMAN	wb52c05.x1 NCI_CGAP_GC6 Hamo sepiens cDNA clone IMAGE:2309286 3' similar to TR:P97213 P97213 CDU2, CDU1, TCDD, TCDB, TCDE, TCDA, TCDC, CDD1, CDD2, CDD3, AND CDD4 GENES.;
7245	17122		1.97	8.0E-71	1171451.1	EST_HUMAN	zp21d11.r1 Stratagene neuroepithellum (#937231) Homo sepiens cDNA clone IMAGE:610101 5' similar to TR:G1143061 G1143061 STRAIN XA34 POL.;
සස	16226	26386	7.91	7.0E-71		EST_HUMAN	zv60h06.r1 Soares_testis_NHT Homo saptens cDNA clone IMAGE:758075 5'
7037	16914		1.52		7.0E-71 AA705457.1	EST_HUMAN	zj91a06.s1 Soares_fetal_liver_spleen_1NFLS_S1 Homo saplens cDNA clone IMAGE:462226 3'
8643	18507	28786	4.18			ΝΤ	Homo sapiens chromosome 21 segment HS21,0010
2163	12050	21951	3.45			L'A	Homo sapiens SP100-HMG nuclear autoantigen (SP100) mRNA, complete cds
4030	13933	23710	1.38		V816405.1	EST_HUMAN	QV4-ST0234-181199-037-f05 ST0234 Homo saplens cDNA
5144	15011	24782	3.2	5.0E-71 AI	829496.1	EST_HUMAN	w/18h10.x1 NCI_CGAP_Ut1 Homo sapiens cDNA clane IMAGE:2425315 3
9855	15501	25577			4502740	LN	Homo sapiens cyclin-dependent kinase 6 (CDK6) mRNA
6434	16295		1.59			NT	Human neurofibromatosis protein type 1 mRNA, 3' end of cds
6548	18408	26585	19.78		5.0E-71 AF072810.1	NT	Homo sapiens transcription factor WSTF mRNA, complete cds
7702	17552		226			MT	Human PreA4 gene for Alzheimer's disease A4 amyloid protein precursor (exon 2)
83.48	18228	28477	10		11436514 NT	Ę	Homo sapiens pro-platelet basic protein (includies platelet basic protein, beta-thromboglobulin, connective tissus-activating paptide-2) (PPBP), mRNA
8528	18400					IN	Homo sapiens similar to hypothetical protein FLU20163 (H. saplens) (LOC63325), mRNA
2906	18848		1.84	L	11417862 NT	NT	Homo saplens calcineurin binding protein 1 (KiAA0330), mRNA
9411	19063		1.62		11418039 NT	NT	Homo sapiens RNA binding motif protein 9 (RBM9), mRNA
97	10082	19899	1.13	4.0E-71	4507592 NT	NT	Homo sapiens tumor necrosis factor (ligand) superfamily, member 10 (TNFSF10) mRNA
347	10306	20123	115.63	4.0E-71 AF	AF157626.1	TN	Equus caballus glyceraldehyde-3-phosphate dehydrogenase mRNA, partial cds
347	10306	20124	115.63	4.0E-71 A	-15762	Ę	Equus caballus glyceraldehyde-3-phosphate dehydrogenase mRNA, partial cds
2850	12778		0.88	4.0E-71	7705414 NT	۲	Homo sapiens hook1 protein (HOOK1), mRNA,
2850	12778					N	Homo sapiens hook1 protein (HOOK1), mRNA
2857					_	۲	Homo sapiens plasminogen (PLG) mKNA
4330					-05632	님	Homo saplens SP100-HMG nuclear autoantigen (SP100) mRNA, complete cds
4913	14792	24567	4.99	4.0E-71	7657602 NT	NT	Homo sapiens putative heme-binding protein (SOUL), mKNA

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Top Hit Descriptor	nI45h10.s1 NCI_CGAP_Pr4 Homo sepiens cDNA clone IMAGE:1043683 similar to contains PTR5.t3 PTR5 repetitive element ;	Homo sapiens chromosome 21 segment HS21,0006	Human mRNA for KIAA0272 gene, partial cds	Human mRNA for KIAA0272 gene, partial cds	Homo sapiens short chain L-3-hydroxyacy/-Co ^λ , dehydrogenase precursor (HADHSC) gene, nuclear gene encoding mitochondrial protein, complete cds	Homo saplens short chain L-3-hydroxyacyl-Cork dehydrogenase precursor (HADHSC) gene, nuclear gene encoding mitochondrial protein, complete cds	bb81a08.y1 NIH_MGC_10 Homo septiens cDNA clone IMACE:3048784 5' similar to SW:R23B_HUMAN P64727 UV EXCISION REPAIR PROTEIN PROTEIN RAD23 HOMOLOG B:	ye43e09.r1 Soares fetal liver spleen 1NFLS Homo saplens cDNA clone IMAGE:120520 5	oy15e03.s1 Soares, senescent fibroblasts, NbHSF Homo sapiens cDNA clone IMAGE:1665916 3' similar to contains LOR1.b2 LOR1 repetitive element;	Homo sapiens neuronal cell death-related protein (LOC51616), mRNA	Homo saplens disabled-2 gene, exons 2 through 15 and complete cds	Homo sapiens phosphatidylinositol 4-kinase 230 (pi4K230) mRNA, complete cds	Homo sapiens PMS2L16 mRNA, partial cds	Homo sapiens PMS2L16 mRNA, partial cds	Homo sapiens hairy/enhancer-of-split related with YRPW motif-like (HEYL), mRNA	Homo sapiens inorganic pyrophosphatase mRNA, complete cds	Homo saplens SNARE protein kinase SNAK mRNA, complete cds	Homo sapiens SNARE protein kinase SNAK mRNA, complete cds	02_15 Human Epidermal Keratinocyte Subtraction Library- Upregulated Transcripts Homo sapiens cDNA done 02_15 5' similar to Homo sapiens chromosome 19	02_15 Human Epidermal Keratinocyte Subtraction Library- Upregulated Transcripts Homo saplens cDNA	Homo sapiens attractin precursor (ATRN) gene expn 19	Human mRNA for KIAA0045 gene, complete cds	Homo sepiens GCN5 (general control of amino-acid synthesis, yeast, homolog)-like 2 (GCN5L2). mRNA	Homo sapiens CAGL 79 mRNA, partial cds	Hamo saplens myomesin (M-protein) 2 (165kD) (MYOM2), mRNA	Homo sapiens hypothetical protein FLJ10998 (FLJ10998), mRNA
Top Hit Database Source	EST_HUMAN	NT	TN	NT	۲	ΙN	EST HUMAN	EST HUMAN	EST HUMAN	NT	NT	N	Z	TN	NT	NT	NT	LN	EST_HUMAN		NT TOWN	N			TN	NT
Top Hit Acession No.	3.0E-71 AA557683.1	2.0E-71 AL163206.2		2.0E-71 D87462.1	F095703.1	F095703.1	2.0E-71 BE018477.1	T95489.1	1.0E-71 AI077927.1	7706281 NT	1.0E-71 AF205890.1			1.0E-71 AB017007.1	57153			1.0E-71 AF246219.1	1.0E-71 BE122850.1	1 05 34 85422850 4		1.0E-71 D28476.1	11426182 NT	U80753.1	1	8922811 NT
Most Similar (Top) Hit BLAST E Value	3.0E-71	2.0E-71	2.0E-71	2.0E-71	2.0E-71 A	2.0E-71 A	2.0E-71	2.0E-71 T	1.0E-71	1.0E-71	1.0E-71	1.0E-71	1.0E-71	1.0E-71	1.05-71	1.0E-71	1.0E-71	1.0E-71	1.0E-71	10 1	1.0E-71	1.0E-71	1.0E-71	1.0E-71	1.0E-71	1.0E-71
Expression Signal	3.32	2:52	6.88	98'9	2.56	2.56	2.3	6.22	2.11	1.83	4.01	10.59	1.23	1.23	4.85	1.17	4.73	4.73	0.94	70 0	1.87	1.86	4:	10.62	6.96	4.18
ORF SEQ ID NO:	28218	89602	24957	24958	28114	281.15	28219		20371	20699	20841	21080	21821	21822	22416	23179	23256	23257	23298	22200	23381	24050	26079	26351	26816	26986
Exan SEQ ID NO:	17968	11119	15181	15181	17872	17872	17970	18920	10559	10851	11000			11927	12528		13462	13462	13511	13511	13595	14268	ļ	ı	ı	16794
Probe SEQ ID NO:	8077	1210	5259	5259	8022	8022	8078	9181	622	926	1084	1317	2036	2036	2661	3467	3546	3546	3597	3507	3681	4370	6044	8326	6750	6916

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Top Hit Descriptor	Homo sapiens hypothetical protein FLJ10998 (FLJ10998), mRNA	Homo sapiens cytochrome c oxidase subunit VIIa-related protein gene, complete cds	AV761217 MDS Homo saplens cDNA clone MDSEIA03 5'	AV781217 MDS Homo sapiens cDNA clone MDSEIA03 5	Homo sapiens coagulation factor XIII, A1 polypeptide (F13A1), mRNA	Homo sapiens leucy//cystiny/ aminopeptidase (LNPEP), mRNA	Homo sapiens leucy//cystiny/ aminopeptidase (LNPEP), mRNA	Homo sapiens gene for AF-6, complete cds	wk95g03.x1 NCI_CGAP_Lu19 Homo septens cDNA clone IMAGE:2423188 3' similar to TR:O86705 O86705 HYPOTHETICAL 38.6 KD PROTEIN: ;contain's Alu repetitive element;	wk95g03.x1 NCI_CGAP_Lu19 Homo sepiens cDNA clone IMAGE:2423188 3' similær to TR:OB6705 O86705 HYPOTHETICAL 38.6 KD PROTEIN. ;contains Alu repetitive element;	Homo sapiens aconitase 2, mitochondrial (AC¢2), nuclear gene encoding mitocondrial protein, mRNA	Homo sapiens aconitase 2, mitochondrial (AC¢2), nuclear gene encoding mitocondrial protein, mRNA	Homo sepiens aconitase 2, mitochondrial (AC¢2), nuclear gene encoding mitocondrial protein, mRNA	(pseudogene) PTMAP2=prothymosin alpha [human, Genomic, 1192 nt, segment 2 of 3]	Homo sapiens chromosome 21 segment HS21 C046	7K63a05.x1 NCI_CGAP_GC6 Homo seplens cDNA clone IMAGE:3480080 3' similar to SW:KMLC_RABIT P07313 MYOSIN LIGHT CHAIN KINASE, SKELETAL MUSCLE;	QV0-CS0010-150900-398-e11 CS0010 Homo sapiens cDNA	QV0-CS0010-150900-398-e11 CS0010 Homo sapiens cDNA	QV0-CS0010-150900-398-e11 CS0010 Homo sapiens cDNA	QV0-CS0010-150900-398-e11 CS0010 Homo sapiens cDNA	Homo sapiens alpha-tubulin mRNA, complete cds	AU128584 NT2RP2 Homo saplens cDNA clone NT2RP2003761 6'	au80c03.y1 Schneider fetal brain 00004 Homo sapiens cDNA clone IMAGE:2782584 5' similar to TR:Q99785 Q99786 HYPOTHETICAL 32.4 KD PROTEIN ; contains element MSR1 repetitive element ;	MR4-BT0598-010600-005-d05 BT0598 Homo sapiens cDNA	MR4-BT0598-010600-005-d05 BT0598 Homo sapiens cDNA	QV1-BT0632-280800-342-a10 BT0632 Hamo saplens cDNA
Top Hit Database Source	ΤN	TN	EST_HUMAN	EST_HUMAN	NT	Z	NT	NT	EST_HUMAN	EST_HUMAN	F	۲Z	NT	L'N	LN L	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	ΝΤ	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN
Top Hit Acession No.	8922811 NT	AY007643.1	AV761217.1	AV761217.1	11418903 NT	11417191 NT	11417191 NT	AB011399.1	Al857635.1	AI857635.1	4501868 NT	4501866 NT	4501868 NT	S41694.1	AL163246.2	BF059578.1	BF333707.1	BF333707.1	BF333707.1	BF333707.1	L11645.1	AU128584.1	AW161274.1	BF331571.1	BF331571.1	BE926645.1
Most Similar (Top) Hit BLAST E Value	1.0E-71	1.0E-71		1.0E-71	1.0E-71	1.0E-71	1.0E-71	1.0E-71	9.0E-72	9.0E-72	7.0E-72	7.0E-72	7.0E-72	7.0E-72	8.0E-72	6.0E-72	5.0E-72	6.0E-72	5.0E-72	5.0E-72	6.0E-72	5.0E-72	5.0E-72	5.0E-72	5.0E-72	5.0E-72
Expression Signal	4.18	6.49	3.39	4.87	3.2	2.33	2.33	4.48	1.15	1.15	6.23	6.23	5.23	2.94	3.72	222	0.88	0.88	2.95	2.95	2.76	1.47	3.55	3.18	3.18	2.43
ORF SEQ ID NO:	26987	27820			28383		28624		20173	20174	23700	23701	23702	26244		28893	19854	19855	19854	19855		26163	27167	28706		
Exan SEQ ID NO:	16794	17598	17642	18052	18136	18359	18359	19147	10347	10347	13928	13926	13926	16094	16784	18603	10043	10043	10043	10043	11037	16023	16974	18437		19700
Probe SEQ ID NO:	6916	7748	7792	8164	8258	8486	8488	9547	401	401	4023	4023	4023	6228	6885	8788	99	99	57	25	1122	9150	7607	8569	8569	9253

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Top Hit Descriptor	Homo sapiens myosin, heavy polypeptide 13, skeletal muscle (MYH13), mRNA	RC4-HT0578-170300-012-g02 HT0578 Homo saplens cDNA	RC4-HT0578-170300-012-g02 HT0578 Homo sapiens cDNA	Homo saplens synaptic glycoprotein SC2 (SC2) mRNA, complete cds	Homo sapiens synaptic glycoprotein SC2 (SC2) mRNA, complete cds	MRo-CT0063-071099-002-h11 CT0063 Homo sapiens cDNA	Homo sapiens ribosomal protein L13a (RPL13A), mRNA	ws55c06.x1 NCI_CGAP_Brn25 Home saplens cDNA clone IMAGE:2501098 3' similar to TR:Q58050 Q58050 HYPOTHETICAL PROTEIN MJ1656.;	ov39h08.x1 Soaree_testis_NHT Homo sapiens;cDNA clone IMAGE:1639743 3'	Homo sapiens lysozyme homolog (LOC57151); mRNA	Homo sapiens vacuolar ATPase Isoform VA68 mRNA, complete cds	bb62a06.y1 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:3030034 5' similar to gb:X04098_cds1 ACTIN, CYTOPI ASMIC 2 (HI MAN): cht.M21405 Marite cytoskeletel gamma-actin mRNA complete cds (MOI ISE)	Homo sapiens interleukin 12 receptor, beta 1 (III.12RB1), mRNA	Homo sapiens Interleukin 12 receptor, beta 1 (III 12RB1), mRNA	Homo sapiens DNA for Human P2XM, complete cds	Homo sapiens thyroid autoantigen 70kD (Ku antigen) (G22P1), mRNA	Homo sapiens hypothetical protein FLJ20309 (FLJ20309), mRNA	Homo saplens chromosome 21 segment HS21(2006	Homo sapiens chromosome 21 segment HS21¢082	Homo sapiens chromosome 21 segment HS210018	QV0-HT0494-020300-137-d03 HT0494 Homo saplens cDNA	Homo sapiens HELG protein (FAM4A1), mRNA	Homo saplens hemo-binding protein (HEBP), mRNA	Homo sapiens heme-binding protein (HEBP), mRNA	Homo sapiens BASS1 (BASS1) mRNA, partial cds	RC3-NN0066-270400-011-c04 NN0066 Homo saplens cDNA	Human beta globin region on chromosome 11	Homo sapiens caspase 8, apoptosis-related cysteine protease (CASP8) mRNA	Homo sapiens Parkinson disease (autosomal recessive, juvanile) 2, parkin (PARK2), transcript variant 3, mRNA
Top Hit Database Source	N N	EST_HUMAN	EST_HUMAN	N	F	EST_HUMAN	INT	EST HUMAN	EST_HUMAN	본	N	NOW I	LN - LN	ĮN.	r _Z	N _T	INT	ħ	TN	NT	EST_HUMAN	INT	INT	TN	IN	EST_HUMAN	NT	INT	TN.
Top Hit Acession No.	11321578 NT	BE175434.1	BE175434.1	AF222742.1	AF222742.1	AW374968.1	11424099 NT	AW071755.1	AI024877.1	11426469 NT	AF113129.1	100000	11526037 NT	11526037 NT	AB002059.1	11418189 NT	8923290 NT	AL163206.2	AL163282.2	AL163218.2	BE166574.1	11422159 NT	11435913 NT	11435913 NT	AF139897.1	AW898081.1	U01317.1	4502582 NT	7669539 NT
Most Similar (Top) Hit BLAST E Value	1.0E-72	1.0E-72	1.0E-72	1.0E-72	1.0E-72	9.0E-73	9.0E-73	8.0E-73	8.0E-73	8.0E-73	8.0E-73	9 0 10	8.0E-73	8.0E-73	8.0E-73	8.0E-73	7.0E-73	7.0E-73	7.0E-73	6.0E-73	6.0E-73	4.0E-73	3.0E-73	3.0E-73	2.0E-73	2.0E-73	2.0E-73	2.0E-73	2.0E-73
Expression Signal	19.78	3.82	3.82	90.9	90.9	1.23	23.9	1.03	3.06	4.6	2	4 0	222	222	2.12	2.69	0.78	1.08	1.62	2.37	3.36	2.05	66.0	66.0	1.75	1.48	1.1	3.48	0.86
ORF SEQ ID NO:	25977	26533		27594	27595			. 20782			26785	077460	27682	27683	25282	26235	20875	22983			26274	24832			20610			22869	23221
Exon SEQ ID NO:	15854	16360	16360	17383	17383	11348	18197	10939	l	15864	i	17264	17465	17465	19081	19217	11033	13184	14743	10126	16121	15138	11715	11716	10760	11798	12135	13069	13420
Probe SEQ ID NO:	5949	6501	6501	7532	7532	1443	8320	1022	1389	5959	6715	7395	7614	7614	9446	9654	1118	3261	4863	152	6255	5215	1818	1818	833	1902	2251	3144	3503

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
3603	13420		0.96	2.0E-73	7669539 NT	NT	Homo sapiens Parkinson disease (eutosomal recessive, juvenile) 2, parkin (PARK2), transcript variant 3, mRNA
5902	15808	25933		2.0E-73 AB	AB046811.1	TN	Homo sapiens mRNA for KIAA1591 protein, partial cds
6023	15927		1.35	2.0E-73	11431471 NT	ΙN	Homo sepiens Interleukin 4 receptor (IL4R), mRNA
6023	15927		1.35	2.0E-73	11431471 NT	ΙN	Homo saplens Interleukin 4 receptor (IL4R), mRNA
7956	17806			2.0E-73	4504168 NT	Ę	Homo sapiens glutathione synthetase (GSS) mRNA
7983	17843				11496980 NT	۲	Homo sapiens supervillin (SVIL), transcript variant 1, mRNA
7993	17843		2.45		11496980 NT	Ę	Homo saplens supervillin (SVIL), transcript variant 1, mRNA
94.39	48028			2 OE-73	11431588 NT	5	Homo sapiens KIAA1080 protein, Golgi-associated, gamme-adaptin ear containing, ARF-binding protein 2 (KIAA1080), mRNA
8408	18284	Ţ		2.0E-73		F	Homo sapiens galactosylceramidase (Krabbe disease) (GALC), mRNA
8408	18284	L		2.0E-73	4557612 NT	۲	Homo sapiens galactosylceramidase (Krabbe disease) (GALC), mRNA
8432	18306	28562		2.0E-73	2.0E-73 AB028982.1	N	Homo saplens mRNA for KIAA1059 protein, pertial cds
9447	11798		1.72	2.0E-73 AV	AW898081.1	EST_HUMAN	RC3-NN0068-270400-011-c04 NN0066 Homo, saplens cDNA
1743	11644			1.0E-73	1.0E-73 AU121585.1	EST_HUMAN	AU121585 MAMMA1 Homo saplens cDNA clone MAMMA1000490 5'
2434	12311	22207		1.0E-73	1.0E-73 AF198349.1	NT	Gallus gallus Dach2 protein (Dach2) mRNA, complete cds
							gg61b07.r1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1839637 5' similar to contains element
7469	17329	27534	1.36	1.0E-73 AI	AI147427.1	EST_HUMAN	MER22 repetitive element ;
8747	17896	28140	2.67	1.0E-73	1.0E-73 BE385477.1	EST_HUMAN	601276071F1 NIH_MGC_20 Homo sapiens cDNA clone IMAGE:3817105 5
723	10655		1.39	8.0E-74	4557426 NT	TN	Homo sapiens CD39-like 4 (CD39L4) mRNA
9099	15521		1.84	8.0E-74	S83194.1	TN	Ca2+/calmodulin-dependent probin kinase IV kinase isoform [rats, brain, mRNA, 3429 nt]
9099			1.84		8.0E-74 S83194.1	TN	Ca2+/calmodulin-dependent protein kinase IV kinase Isoform [rats, brain, mRNA, 3429 nt]
1906			3.01	7.0E-74	7.0E-74 AJ001689.1	NT	Homo sapiens NKG2D gene, exon 10
3286	13207	23007	66.0			NT	Homo sapiens chromosome 21 segment HS21,0046
7327	17231	27432	2	7.0E-74	7.0E-74 BE967432.1	EST_HUMAN	601649284F1 NIH_MGC_73 Hamo saplens cDNA clone IMAGE:3932897 5'
9653	19216	25234	2.81	7.0E-74		EST_HUMAN	601191927F1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:3535855 5'
	l					ŀ	Homo sapiens S164 gene, partial cds; PS1 and hypothetical protein genes, complete cds; and S171 gene,
1106			2.58		AF109907.1	2	Parties Cas
1609	11514	21373	1.03		AW263177.1	EST_HUMAN	xn78g07.x1 Soares_NFL_T_GBC_S1 Homo sepiens cLINA clone IMAGE:2/00636 3
2268	12152				6.0E-74 BE388260.1	EST_HUMAN	601283521F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3605453 5
2268	12152		96.6		6.0E-74 BE388260.1	EST HUMAN	601283521F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3605453 5
2834					6.0E-74 AW014039.1	EST_HUMAN	UI-H-BI0-aah-h-03-0-UI.s1 NCI_CGAP_Sub1 Homo sapiens cDNA clone IMAGE:2709365 3
2834					6.0E-74 AW014039.1	EST_HUMAN	UI-H-BI0-aah-h-03-0-UI.s1 NC CGAP Sub1 Homo sapiens cDNA clone IMAGE:27/09355 3
3652	13566	23352	2.63		6.0E-74 BE048846.1	EST_HUMAN	hr54e11.x1 NCL_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:3132332.3

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Table 4
Single Exon Probes Expressed in Heart

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	Top Hit Descriptor	hr54e11.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:3132332 3'	Homo sapiens actin filament associated protein (AFAP), mRNA	df17c09.y1 Morton Fetal Cochled Homo sepiens cDNA clone IMAGE:2483704 5'	PM0-CT0289-271099-001-h07 CT0289 Homo sapiens cDNA	Homo sapiens phosphatidylinositol glycan, class L (PIGL), mRNA	H.sapiens mRNA for TPCR16 protein	Homo sapiens VAMP (vesicle-associated membrane protein)-associated protein A (33kD) (VAPA) mRNA, and translated protein te	Homo capiens interlegible 4 recentor (II 4P) mRNs	Homo sapiens interlaukin 4 receptor (IL4R), mRNA	Homo sapiens KIAA0716 gene product (KIAA0716), mRNA	Homo saplens hypothetical protein FLJ13222 (FLJ13222), mRNA	H.saplens mRNA for HIP-I	H.sapiens mRNA for HIP-I	Homo saplens DNA for amyloid precursor protein, complete cds	Homo sapiens mRNA for KIAA1019 protein, partial cds	Homo sapiens DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes, complete eds)	Homo sapiens DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes,	complete ods)	Homo sapiens proteasome (prosome, macropain) subunit, beta type, 1 (PSMB1) mRNA	Homo sapiens proteasome (prosome, macropain) subunit, beta type, 1 (PSMB1) mRNA	Homo saplens mRNA for KIAA1168 protein, partial cds	Homo sapiens PLP gene	Homo saplens PLP gene	Homo sapiens chromosome 21 segment HS21C010	Homo saplens chromosome 21 segment HS21C047	Homo sapiens KIAA0569 gene product (KIAA0559), mRNA	Homo sapiens mRNA for transmebrane receptor protein	Homo sapiens hydroxyacyl-Coenzyme A dehydrogenase/3-ketoacyl-Coenzyme A thiolase/enoyl-Coenzyme A hydratase (tritinoclinnel protein). Peta submit (HADHR) mRNA	Homo sapiens hydroxyacy-Coenzyme A dehydrogenase/3-ketoacy/-Coenzyme A thiolase/enoy/-Coenzyme A hydrotase (trifunctional protein), beta subunit (HADHB) mRNA	EST13131 Thymus tumor III Homo sapiens cDNA 5' end similar to similar to ribosomal protein L37
, -	Top Hit Database Source	EST_HUMAN	- LZ	EST_HUMAN	EST_HUMAN	Z	F	·		LZ	Z	۲	NT	TN.	L	FZ	N		LZ.	NT	NT	NT	LN	NT	. LN	NT	F	N T	F	F	EST_HUMAN
	Top Hit Acession No.	BE048846.1	11056013 NT	4W020986.1	4W362756.1	11425417 NT	X89870.1	745078BB	11434474 NT	11431471 NT	7662263 NT	11345483 NT	Y09420.1	r09420.1	J87675.1	AB028942.1	AB026898.1	:	4B026898.1	4506192 NT	6192		A_J006976.1	AJ006976.1	AL163210.2	AL163247.2	7662183 NT	217227.1	TN 902304	4504328 NT	AA300378.1
	Most Similar (Top) Hit BLAST E Value	6.0E-74	6.0E-74	5.0E-74	5.0E-74	5.0E-74	5.0E-74	5.0E_74	A7-70.2	5.0E-74	5.0E-74	5.0E-74	6.0E-74	5.0E-74	4.0E-74	4.0E-74	4.0E-74		4.0E-74 /	4.0E-74	4.0E-74	4.0E-74	4.0E-74	4.0E-74	4.0E-74	4.0E-74	4.0E-74	4.0E-74	4 0F-74	4.0E-74	3.0E-74
	Expression Signal	2.63	2.49	2.58	5.19	2.15	10.48	A 7.4	1 85	1,85	3.73	2.69	1.88	1.88	1.89	4.95	2.44		2.44	4.34	4.34	1.21	0.99	4.44	0.83	1.31	1.57	0.82	9.28	3.76	5.03
	ORF SEQ ID NO:		25016					2555				26758	28252	28253	20063	20811	21692							22772	23204	23655	24142	24192	24844		Ш
	Exon SEQ ID NO:	13566	L	10813	12534	15240	15456	15479	15518	15516	16014	16564	18006	18006	10243	10761	11814		11814	11918	11918	11975	12257	12979	13399	13880	14351	14405	14880	j	, ,
	Probe SEQ ID NO:	3652	5294	887	2669	5320	9239	5563	5802	5602	6120	6884	8117	8117	278	834	1919		1919	2027	2027	2085	2377	3052	3483	3973	4457	4512	5006	5008	6977

WO 01/57274 PCT/US01/00666

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Top Hit Descriptor	EST01132 Subtracted Hippocampus, Stratagene (cat. #936205) Homo sapiens cDNA clone HHCPF91	no17g05.s1 NCI_CGAP_Phe1 Hamo sapiens cDNA clone IMAGE:1100984 3'	Homo sapiens glyceraldehyde-3-phosphate dehydrogenase (GAPD), mRNA	Homo sapiens glyceraldehyde-3-phosphate dehydrogenase (GAPD), mRNA	Human endogenous retrovirus HERV-K-T47D	wx51e07.x1 NCI_CGAP_Lu28 Homo septens cDNA done IMAGE:2547204 3' similar to SW.GG95_HUMAN Q08379 GOLGIN-95, contains element MER22 repetitive element;	Homo saptens epidermal growth factor receptor (avian erythrobiastic leukemia viral (v-erb-b) oncogene homolog) (EGFR) mRNA	Homo sapiens epidermal growth factor receptor (avian erythroblastic leukemia viral (v-erb-b) oncogene	Inditional (EGTK) MINNA	P.1.2.113_c11.r tumorz Homo sapiens CLNNA 3	Nover numan gene mapping to cnomosome 22	Ivova naman gene mapping to comosome zz	Human pietelet glycoprotein ilb mKNA, 3' end		nomo sapiens ruc-ra protein (Puz-rain r-Cu-38), minna	Home sapiens PUZ-73 promein (PUZ-73/NY-CU-38), mKNA	Home saplens PDZ-73 protein (PDZ-73/NY-CC-38), mKNA	Homo sapiens PDZ-73 protein (PDZ-73/NY-CQ-38), mRNA	601557524F1 NIH_MGC_58 Homo sapiens cDNA clone IMAGE:3827549 5'	Homo saplens mRNA for KIAA1395 protein, partial cds	Homo saplens chromosome 21 segment HS210004	др96a06.s1 Stratagene muscle 937209 Homo sapiens cDNA clone IMAGE:628018 3'	602121428F1 NIH_MGC_56 Homo saplens cDNA clone IMAGE:4278559 6'	Homo sapiens Misshapen/NIK-related kinase (MINK), mRNA	QV4-ST0234-181199-037-105 ST0234 Homo saplens cDNA	Homo saplens hypothetical protein FLJ11026 (FLJ11026), mRNA	Homo saplens beta 2 gene	Homo saplens zinc finger protein 259 (ZNF259) mRNA	Homo sapiens chromosome 21 segment HS21C046	Homo sapiens DNA for Human P2XM, complete cds	Homo saplens mannosidase, alpha, class 2A, member 1 (MAN2A1), mRNA
Top Hit Database Source	EST_HUMAN	EST_HUMAN	Z	Σ	FX	EST HUMAN			ıŀ	EST HUMAN	z	IN I	TANK TANK	-1	Ž	- 1	Z	Z	EST_HUMAN	NT	LZ L	EST_HUMAN	EST_HUMAN	FN	EST_HUMAN	닐	M	FA	NT	NT	NT
Top Hit Acession No.	M78984.1	AA601493.1	7669491 NT	7669491 NT	AF020092.1	AI950528.1	7885198 NT		20180			ALSSSUSZ. 1	J02963.1	44.400507	N /8000411	11439587 N	11439587 NI	11439587 NT					BF666568.1	7657334 NT	AW816405.1	8922829 NT	X02344.1	4508020 NT		AB002059.1	4758697 NT
Most Similar (Top) Hit BLAST E Value	3.0E-74	3.0E-74	2.0E-74	2.0E-74	2.0E-74	2.0E-74	2.0E-74		2.0E-74	2.0E-/4	2.0E-/4	Z.UE-/4	2.0E-74	2.0E-/4	2.0E-74	2.0E-/4	2.0E-74	2.0E-74	2.0E-74	2.0E-74	2.0E-74	2.0E-74	2.0E-74	1.0E-74	1.0E-74	1.0E-74	1.0E-74	1.0E-74	1.0E-74	1.0E-74	1.0E-74
Expression Signal	2.42	2.22	126.24	126.24	1.01	1.15	2.94		5 8	1.09	08:1	<u> </u>	1.85	5 5	3 5	200	7:2/	2.57	1.55	1.43	6.54	1.46	1.26	76.0	3.6	1.19	2.7	1.35	2.17	6.19	5.96
ORF SEQ ID NO:	27519	28010		20715	20916	20986	21340	3	14017				24588								27523		25198	19841	20108	20247	20252	20331	20751		22822
Exon SEQ ID NO:	17312	17771	10867	10867	11071	11132	11481		10401	┸	14610	┸	14820	┸	Ŀ	⊥			_ [- [- 1		19380			10434	10439	10524	10906		13028
Probe SEQ ID NO:	7394	7921	942	942	1158	1224	1577	1	2660	BCC7	4957	3	4942	2	900	9500	200	5629	6219	6831	7398	9387	9903	47	334	491	. 497	586	983	2179	3100

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Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
1795	5 11693	21569	1.98	3.0E-75	AB011153.1	NT	Homo saplens mRNA for KIAA0581 protein, partial cds
2065	11955		1.05	3.0E-75	4507334 NT	NT	Homo sapiens synaptojanin 1 (SYNJ1), mRNA
2373	3 12253	22144	3.66	3.0E-75	4769153 NT	NT	Homo sapiens synaptosomal-associated protein, 23kD (SNAP29) mRNA
2888	L		1.19	3.0E-75	AL163201.2	NT	Homo sapiens chromosome 21 segment HS21C001
3152		ľ	1.33	3.0E-75	AB011153.1	NT .	Homo saplens mRNA for KIAA0581 protein, partial cds
3306	L	23030	1.01	3.0E-75	M72393.1	NT	Human calcium-dependent phospholipid-binding protein (PLA2) mRNA, complete cds
3306			1.01	3.0E-75	M72393.1	NT	Human calcium-dependent phospholipid-binding protein (PLA2) mRNA, complete cds
4077	13979		1.42	3.0E-75	D87675.1	NT	Homo saplens DNA for amyloid precursor protein, complete cds
4336	14233	24015	96.0	3.0E-75	7662421 NT	TN	Homo sapiens KIAA0971 protein (KIAA0971), mRNA
6058	3 16041	26183	1.53	3.0E-75	11526319 NT	TN	Homo sapiens HIR (histone cell cycle regulation defeative, S. cerevislae) homolog A (HIRA), mRNA
6058			1.53	3.0E-75	11528319 NT	۲N	Homo sapiens HIR (histone cell cycle regulation defective, S. cerevisiae) homolog A (HIRA), mRNA
6234		_	3.83	3.0E-75	7662209 NT	TN	Homo sapiens KIAA0623 gene product (KIAA0623), mRNA
6234				3.0E-75		LN	Homo saplens KIAA0623 gene product (KIAA0623), mRNA
6494	1		2.82	3.0E-75	4885632 NT	NT	Homo saplens Oncogene TIM (TIM) mRNA,
6494	4 16353	26523	2.82	3.0E-75	4885632 NT	NT	Homo sapiens Oncogene TIM (TIM) mRNA
7193	L		1.23	3.0E-75	11420804 NT	IN	Homo sapiens snail 1 (drosophila homolog), zinc finger protein (SNA11), mRNA
5480	15400		1.5	2.0E-75	AV734680.1	EST_HUMAN	AV734680 cdA Homo sapiens cDNA clone cdABED02 5
7082		27152	1.73	2.0E-75	Al311783.1	EST HUMAN	qo91e02x1 NCI_CGAP_Kid5 Homo sapiens cDNA clone IMAGE:1915898 3' similar to TR:Q69386 Q69386 POL/ENV GENE;
							xg60d02.x1 NCI_CGAP_Ut4 Homo sapiens cDNA clone IMAGE:2632707 3' similar to contains PTR7.t1
2255		22037	5.09	1.0E-75	AW168135.1	EST HUMAN	PTR7 repetitive element;
2916	12843		3.17	1.0E-75	X52221.1	NT	H.sapiens ERCC2 gene, exons 1 & 2 (partial)
							2457h03.s1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:726485 3' similar to gb:M13932 40S
6903					AA399270.1	EST_HUMAN	RIBOSOMAL PROTEIN S17 (HUMAN);
7423	3 17290	27499	3.73	1.0E-76	BF313645.1	EST_HUMAN	601900294F1 NIH_MGC_19 Home septems cDNA clare IMAGE:4128678 5
7423	3 17290	27500	3.73	1.0E-75	BF313645.1	EST_HUMAN	601900294F1 NIH_MGC_19 Homo sapiens cDNA clane IMAGE:4129678 5
8257	7 18137		3.90	1.0E-76	AA664377.1	EST_HUMAN	ac77b08.s1 Stratagene lung (#937210) Homo sépiens cDNA clone IMAGE:868599 3'
							Homo sapiens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively
8441	1 18315	28573	2.6	1.0E-75	AF223391.1	NT	spliced
9299	15100	24892	1.86	1.0E-75	BE894192.1	EST_HUMAN	601437130F1 NIH_MGC_72 Homo sapiens cDNA done IMAGE:3922303 5
gg	10026	19875	1.45	9.0E-76	Al652648.1	EST HUMAN	wb30b10.x1 NCI_CGAP_GC6 Homo seplens cDNA clone IMAGE:2307163 3' similar to TR:O75235 O75235 TRAP1 ;
	1	١					

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Top Hit Descriptor	wb30b10.x1 NCI_CGAP_GC8 Hamo sepiens cDNA clone IMAGE:2307163 3' similar to TR:075235 075235 TRAP1;	z183b07.s1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:447541 3'	Human ferritin Heavy subunit mRNA, complete cds	Homo sapiens H factor 1 (complement) (HF1) inRNA	Homo sapiens H factor 1 (complement) (HF1) inRNA	Homo sapiens mediator (Sur2), mRNA	Homo sapiens LIM domain kinase 1 (LIMK1), niRNA	Homo saplens serine/threonine kinase 2 (STK2), mRNA	Homo sapiens baculoviral IAP repeat-containing 6 (BIRC6), mRNA	Homo sapiens calcineurin binding protein 1 (KIAA0330), mRNA	Homo sapiens dihydrolipoamide dehydrogenase (E3 component of pyruvate dehydrogenase complex, 2-oxo-	glutarate complex, branched chain keto acid dehydrogenase complex) (ULD) mKNA	Homo sapiens cAMP-specific phosphodiesterase 8A (PDE8A) mRNA, partial cds	Homo sapiens lymphocyte antigen 75 (LY75) mRNA, and translated products	Homo sepiens core-binding factor, runt domain; alpha subunit 2; translocated to, 1; cyclin D-related (OBFA2T1) mRNA	Homo saplens sepiapterin reductase (7,8-dihydrobiopterin:NADP+ oxidoreductase) (SPR) mRNA	Homo sapiens sepiaptenn reductase (7,8-dihydroblopterin:NADP+ oxidoreductase) (SPR) mRNA	601312019F1 NIH_MGC_44 Hamo sapiens cDNA clone IMAGE:3658757 5'	601142253F1 NIH_MGC_14 Hamo saplens cDNA clone IMAGE:3508029 5'	Human mRNA for HMG-1, complete cds	Human mRNA for HMG-1, complete cds	Human mRNA for HMG-1, complete cds	HUM178G01B Human fetal brain (TFujiwara) Homo sapiens cDNA clone GEN-178G01 5'	HUM178G01B Human fetal brain (TFujiwara) Homo saplens cDNA done GEN-178G01 5'	UI-H-BW 1-enz-b-04-0-UI.s1 NCI_CGAP_Sub7, Homo sapiens cDNA clone IMAGE;3083862 3'	UI-H-BW 1-anz-b-04-0-UI,s1 NCI_CGAP_Sub7, Homo sapiens cDNA clone IMAGE:3083862 3'	Homo sapiens eukaryotic translation elongation factor 1 beta 2 (EEF1B2) mRNA	Homo sapiens eukaryotic translation elongation factor 1 beta 2 (EEF1B2) mRNA	RC5-ST0300-180100-033-A03 ST0300 Homo saplens cDNA	RC5-ST0300-180100-033-A03 ST0300 Homo capiens cDNA	HSCZQD042 normalized Infant brain cDNA Homo sapiens cDNA clone c-zqd04 3'
Top Hit Database Source	EST_HUMAN	EST_HUMAN	LZ LZ															EST_HUMAN	EST_HUMAN	NT		ĹΝ	EST_HUMAN		EST_HUMAN	EST_HUMAN			EST_HUMAN		EST_HUMAN
Top Hil Acession No.	9.0E-76 AI652648.1	9.0E-76 AA702415.1		4504374 NT	4504374 NT	7706724 NT	11421442 NT	11435215 NT	10442821 NT	11417862 NT		5016092	F058490.1	4505052 NT	4757915 NT	4507184 NT	4507184 NT	E396253.1	E273201.1					,	2.1	F516262.1	4503476 NT	4503476 NT	F375689.1	F375689.1	241314.1
Most Similar (Top) Hit BLAST E Value	9.0E-76	9.0E-76	9.0E-76 M12937.1	8.0E-76	8.0E-76	8.0E-76	8.0E-76	8.0E-76	8.0E-76	8.0E-76		7.0E-76	7.0E-78 A	7.0E-76	7.0E-76	7.0E-76	7.0E-76	6.0E-76	6.0E-76	6.0E-76 D63874.1	5.0E-76 D63874.1	5.0E-76 D63874.1	4.0E-76 D81625.1	4.0E-76 D81625.1	3.0E-76	3.0E-76	3.0E-76	3.0E-76	3.0E-76	3.0E-76	3.0E-76 Z41314.1
Expression Signal	1.45	1.16	23.8	1.69	1.69	1.08	5.38	1.3	6.44	1.44		4.12	2.64	7.08	7	4.3	4.3	19.65	2.69	3.69	3.69	3.69	5.6	5.6	1.78	1.78	2.78	2.78	4.18	4.18	9.13
ORF SEQ ID NO:	19826		27770	20695	20698	22603	25760	26443	28198				22975	28622	23017	l			28154	21673		21675	27834		20359	20380	21345	21346	23089		29104
Exan SEQ ID NO:	10026	12244	17647	10848	10848	<u>_</u>	15652	16282	17948	19207			13177	13183	13215	L		l	17910	11795		11795	17609	17609	10549	10549	11485	11485	13300	l	1 1
Probe SEQ ID NO:	38	2364	7697	923	923	2880	5744	6421	8057	9836		769	3254	3260	3283	4275	4275	1214	8761	1899	1899	1899	7759	7759	613	613	1581	1581	3382	3382	5204

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des journesses	homolog) (PSMD7)			3' similar to contains		RNA	RNA			33'						28 5'	!		Almep			•				190			- Speed Service		
Top Hit Descriptor	Homo sapiens proteasome (prosome, macropain) 26S subunit, non-ATPase, 7 (Mov34 homolog) (PSMD7) mRNA	ze62e02.r1 Soares retina N2b4HR Homo sapians cDNA clone IMAGE:363578 5'	ze62e02.r1 Scares retina N2b4HR Homo capiens cDNA clone IMAGE:363578 5'	ye69f04.s1 Soares fetal liver spleen 1NFLS Homo saplens cDNA clone IMAGE:123007.3 similar to contains MER10 repetitive element:	zu 91 g01.s1 Scares_testis NHT Homo sapiens cDNA clone IMAGE:745392 3'	Homo sapiens polymerase (RNA) II (DNA directed) polypeptide E (25kD) (POLR2E) mRNA	Homo sapiens polymerase (RNA) II (DNA directed) polypeptide E (25kD) (POLR2E) mRNA	Homo sapiens interferon (alpha, beta and omega) receptor 2 (IFNAR2) mRNA	EST369823 MAGE resequences, MAGE Homo sapiens cDNA	qe77h12.x1 Soares_fetal_lung_NbHL19W Horrio sapiens cDNA clone IMAGE:1745063 3'	7 Homo sapiens glucokinase (GCK) gene, exon 2	Homo sapiens disintegrin and metalloprotease clomain 10 (ADAM10) mRNA	Homo sapiens cullin 1 (CUL1) mRNA	Homo sapiens EGF-like repeats and discoidin Iike domains 3 (EDIL3), mRNA	Homo sapiens EGF-like repeats and discoidin I-like domains 3 (EDIL3), mRNA	DKFZp434G1728_r1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434G1728 5	Homo sapiens 3-hydroxylsobutyryl-Coenzyme A'hydrolase (HIBCH), mRNA	Homo saplens 3-hydroxylsobutyryl-Coenzyme Alhydrolase (HIBCH), mRNA	Homo sapiens sorting nextn 5 (SNX5), mRNA	Homo saplens sorting nexin 5 (SNX5), mRNA	Human mRNA for KIAA0299 gene, partial cds	Human mRNA for KIAA0299 gene, partial cds	Homo saplens SET domain and mariner transposase fusion gene (SETMAR) mRNA	Homo saplens SET domain and mariner transposase fusion gene (SETMAR) mRNA	PM3-MT0078-080800-005-g03 MT0078 Homo sapiens cDNA	AV764617 MDS Homo sapiens cDNA clone MDSBTF10 5'	RC3-BN0053-170200-011-h01 BN0053 Homo eaplens cDNA	Homo sapiens CYP17 gene, 5' end	Homo saplens CGI-79 protein (LOC51634), mRNA	Homo sapiens mRNA for KIAA1415 protein, partial cds	
Top Hit Database Source		EST_HUMAN Ze	EST_HUMAN ZE	ST HUMAN	Т				EST_HUMAN E	EST_HUMAN qe						EST_HUMAN DI									EST_HUMAN PI	EST_HUMAN A	EST_HUMAN R				
Top Hit Acession No.	4506230 NT	A019770.1	A019770.1			5944	4505944 NT	4504600 NT	W957753.1	1204066.1	AF041015.1	4567250 NT	4503160 NT	5031660 NT	5031660 NT	5.0E-77 AL043953.1	11428849 NT	11428849 NT	11421928 NT	11421928 NT	AB002297.1	AB002297.1	5730038 NT	5730038 NT		2.0E-77 AV764617.1		-41825.1	7706315 NT	B037836.1	
Most Similar (Top) Hit BLAST E Value	8.0E-77	8.0E-77	8.0E-77	8.0E-77 R00245.1	7.0E-77	7.0E-77	7.0E-77	6.0E-77	6.0E-77	6.0E-77	5.0E-77	6.0E-77	5.0E-77	5.0E-77	6.0E-77	5.0E-77	5.0E-77	6.0E-77	5.0E-77	5.0E-77	. 5.0E-77	5.0E-77	3.0E-77	3.0E-77	3.0E-77 B	2.0E-77	2.0E-77	2.0E-77	2.0E-77	2.0E-77 A	
Expression Signal	2.46	2.12	2.12	7.25	3.28	2.1	2.1	4.29	6.0	17.64	1.78	1.16	0.97	2.02	2.02	2.05	1.39	1.39	2.55	2.55	1.96	1.96	1.12	1.12	3.31	1.71	1.73	0.84	2.64	2.02	
ORF SEQ ID NO:	25095		28862	25232	21659	22136	22137	20043	20882	21287	20973	21101	22492	24280	24281	24514	26947	26948	27543	27544	28080	28081	21705	21706	28377	21093	21185			22312	
Exch SEQ ID NO:	15268	18578	18578	19289	11783	12240							12598	14493	H	14733			1					11824	18129					12724	
Probe SEQ ID NO:	5347	8691	8691	1.22	1887	2360	2360	262	1125	1524	1216	1337	2736	4605	4605	4853	6873	6873	7519	7519	7890	7990	1929	1929	8249	1330	1414	2044	2056	2549	

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Top Hit Database Source	ho43b05x1 Sogres, NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:3040113 3' similar to EST_HUMAN SW:GAG2_HUMAN P10264 RETROVIRUS-RELATED GAG POLYPROTEIN;	tw22g02.x1 NCI_CGAP_Bm52 Homo saplens cDNA clone IMAGE:2260466 3' similar to TR:065245 EST_HUMAN 065245 F21E10.7 PROTEIN ;	tw22g02.x1 NCI_CGAP_Bm52 Homo saplens cDNA clone IMAGE:2260466 3' similar to TR:065245 EST_HUMAN 065245 F21E10.7 PROTEIN	Homo sapiens glutamic-oxaloacetic transaminase 2, mitochondrial (aspartate aminotransferase 2) (GOT2), nuclear gene encoding mitochondrial protein, mRNA	ns68g12.s1 NCI_CGAP_Pr2 Homo sapiens cDNA done IMAGE:1188338 similar to SW:RL29_HUMAN	EST_HUMAN 601119852F1 NIH_MGC_17 Homo sepiens cDNA clone IMAGE:3029436 5'	EST_HUMAN 601476802F1 NIH_MGC_68 Homo sapiens cDNA clone IMAGE:3879505 5'	at74a09.x1 Barstead colon HPLRB7 Homo sapjens cDNA clone IMAGE:2377720 3' similar to TR:Q13311 EST_HUMAN Q13311 TAX1-BINDING PROTEIN TXBP151.[1];	Human protein kinase C substrate 80K-H (PRKCSH) gene, exon 7	Human protein kinase C substrate 80K-H (PRKCSH) gene, exon 7	Homo sapiens mRNA for KIAA1276 protein, partial cds	Homo sapiens mRNA for KIAA1276 protein, partial cds	Homo capiens amyloid beta (A4) precursor protein (protease nexin-II, Alzheimer disease) (APP), mRNA	Homo sapiens amyloid beta (A4) precursor protein (protease nexin-II, Alzheimer disease) (APP), mRNA	Homo sapiens amylold beta (A4) precursor protein (protease nexin-II, Alzheimer disease) (APP), mRNA	Homo sapiens amyloid beta (A4) precursor protiżin (protease nexin-II, Alzheimer disease) (APP), mRNA	Homo sapiens mRNA for KIAA1101 protein, complete cds	Homo sapiens 2,4-dienoy CoA reductase 1, mitochondrial (DECR1), mRNA	Homo sapiens CGI-60 protein (LOC51626), mRNA	Homo sapiens 959 kb contig between AML1 and CBR1 on chromosome 21q22; segment 1/3	Homo sapiens breast cancer 1, early onset (BRCA1), transcript variant BRCA1-exon4, mRNA	CMYA5 Human cardiac muscle expression library Homo saplens cDNA clone 4151935 similar to CMYA5 EST HUMAN Cardiamyopathy associated gene 5
Top Hit Acession No.	2.0E-77 BE044316.1 ES	613519.1	613519.1	4504068 NT	2.0E-77 AA653025.1 ES		2.0E-77 BE787143.1 ES	833003.1	2.0E-77 U50321.1 NT	U50321.1 NT	1.0E-77 AB033102.1 NT	4B033102.1 NT	4502166 NT	4502168 NT	4502168 NT	4502166 NT	AB029024.1 NT	4503300 NT	7706299 NT	4J229041.1 NT	6552322 NT	4 OE 27 AM756264 4
Most Similar (Top) Hit BLAST E Value	2.0E-77	2.0E-77 AI	2.0E-77 AI	2.0E-77	2.0E-77	2.0E-77	2.0E-77	2.0E-77 AI	2.0E-77	2.0E-77	1.0E-77	1.0E-77	1.0E-77	1.0E-77	1.0E-77	1.0E-77	1.0E-77	1.0E-77	1.0E-77	1.0E-77 A	1.0E-77	1 0E-77
Expression Signal	8.1	0.89	0.89	3.48	3.58	1.8	1.34	12.74	4.89	4.99	76.0	26.0	1.87	1.87	4.95	4.95	1.22	2.82	2.99	14.73	1.95	2 89
ORF SEQ ID NO:	23629	23995			24334	25629	25761	26276	27564	27585	19821	19822	20054	20055	20635	20636						24810
Exan SEQ ID NO:	13855		14212	14386	14545	15540	1	16123	17359	17359	10024	10024	10237	10237	12679	12679	12272	12935			14445	15043
Probe SEQ ID NO:	3947	4316	4315	4492	4659	5925	5745	6257	7489	7489	26	4 E	271	271	857	258	2394	3007	4256	4423	4552	6179

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	Top Hit Descriptor	Homo saplens dynactin 1 (DCTN1) gene, exons 27 and 28	Homo sapiens dynactin 1 (DCTN1) gene, exons 27 and 28	Human von Willebrand factor gene, exon 20	Homo sapiens elastin (supravalvular aortio steriosis, Williams-Beuren syndrome) (ELN), mRNA	Homo sapiens hu-GlcAT-P mRNA for glucurory/transferase, complete cds	Homo sapiens hu-GlcAT-P mRNA for glucuroriyitransferase, complete cds	RC3-CT0254-280999-011-b05 CT0254 Homo sapiens cDNA	RC2-ET0023-080500-012-e05 ET0023 Homo sapiens cDNA	RC2-ET0023-080500-012-e05 ET0023 Hamo saplens cDNA	AU118789 HEMBA1 Homo sapiens cDNA clone HEMBA1004354 5'	AU118789 HEMBA1 Homo sepiens cDNA clone HEMBA1004354 5'	Homo saplens GDNF family receptor alpha 1 (GFRA1), mRNA	Homo sapiens hypothetical protein FLJ11316 (FLJ11316), mRNA	ba54h03.y3 NIH_MGC_10 Homo sapiens cDNA clone IMAGE:2900405 5' similar to WP:Y48B6A.6	CE22121;	Human collagenase type IV (CLG4) gene, exon 6	Homo sapiens Best's macular dystrophy related protein mRNA, partial cds	Homo saplens transforming growth factor, beta-induced, 88kD (TGFBI), mRNA	EST365190 MAGE resequences, MAGB Homo sapiens cDNA	Human lysosomal alpha-mannosidase (manB) gene, exon 7	601648061F1 NIH_MGC_62 Homo sapiens cDNA clone IMAGE:3931887 5'	Novel human gene mapping to chomosome 22	wr97b12.x1 NCI_CGAP_Kid11 Homo sapiens ¿DNA clone IMAGE:2495615 3' similar to SW:WAP_PIG 046655 WHEY ACIDIC PROTEIN PRECURSOR;	Homo sapiens pre-mRNA splicing factor (SFRS3) mRNA, camplete cds	Homo sapiens syncytin (LOC30816), mRNA	Homo sapiens phosphatidylinositol 4-kinase, cetalytic, alpha polypeptide (PIK4CA) mRNA	Homo sapiens phosphatidylinositol 4-kinase, catalytic, alpha polypeptide (PIK4CA) mRNA	Homo sapiens hypothetical C2H2 zinc finger protein FLJ22504 (FLJ22504), mRNA	Homo sapiens hypothetical C2H2 zinc finger protein FLJ22504 (FLJ22504), mRNA	Homo sapiens s-CaBP1 (CABP1) mRNA, com slete cds	Human transforming growth factor-beta precursor gene exons 4-5 (and Joined mature peptide)	Homo saplens gene for AF-6, complete cds	Homo sapiens eRF1 gene, complete cds
-	Top Hit Database Source	H	H	H		NT H	H	EST_HUMAN R			EST_HUMAN A	EST_HUMAN A	NT TN			EST_HUMAN C		H		T_HUMAN	드	L'HUMAN	LN	EST HUMAN O	Ì									¥.
	Top Hit Acesslon No.	AF086944.1	AF086944.1	M26844.1	5881412 NT	AB029396.1		AW753302.1	AW947061.1	AW947061.1	AU118789.1	AU118789.1	32710	11422486 NT		AW673424.1	M55586.1	AF038536.1	6585	AW953120.1	U60889.1	BE960836.1	AL355841.1	AI985094.1		7656876 NT	4505806 NT	4505806 NT	11560151 NT	11560151 NT	AF169148.1			AF095901.1
	Most Similar (Top) Hit BLAST E Value	1.0E-77	1.0E-77		1.0E-77		1.0E-77	9.0E-78	8.0E-78	8.0E-78	6.0E-78	6.0E-78	8.0E-78	5.0E-78		5.0E-78	5.0E-78	5.0E-78	5.0E-78	5.0E-78	5.0E-78	5.0E-78	4.0E-78	4.0E-78		4.0E-78	4.0E-78	4.0E-78	4.0E-78	4.0E-78	4.0E-78	4.0E-78	4.0E-78	3.0E-78
	Expression Signal	1.93	1.93	1.66	11.39	1.22	122	2.63	2.97	2.97	1.83	1.83	2.51	1.1		4.1	3.81	2.39	9.35	2.17	6.5	3.75	1.6	1.53	2.21	1.30	1.27	1.27	1.94	1.94	1.97	3	2.85	1.6
	ORF SEQ ID NO:	25607	25608	25698	26089	28093	L	28109	25907		19881	19882		19996		22281	23066	25048						21393					L	28059			Ш	18944
	Exon SEQ ID NO:	15525	15525	15597	15957	17852	17852	17863	15786	16788	10084	10064	1	L		12389	ı		15343	16110	17125	17128	11408	L	L	L	14550	14550		17817	18538		L	10129
	Probe SEQ ID NO:	5610	5610	2688	6197	8002	8002	8013	2880	6880	8	8	9990	211		2515	3339	5323	5422	6244	7248	7249	1502	1628	2270	4227	4684	4684	7867	7967	8721	8844	9664	166

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					>		
Probe E SEQ ID SE NO:	SEQ ID ORF	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
155	10128	19945	1.6	3.0E-78 A	F095901.1	NT	Homo sapiens eRF1 gene, complete cds
3178	13103	22908	0.91	3.0E-78	4507164 NT	NT	Homo sapiens nuclear antigen Sp100 (SP100) mRNA
L	L	23439	0.93	3.0E-78	4507334 NT		Homo saplens synaptojanin 1 (SYNJ1), mRNĄ
,	17762		5.58	3.0E-78	E144758.1	EST_HUMAN	CM0-HT0180-041099-065-c07 HT0180 Homo saplens cDNA
8349	18226	28478	5.52	3.0E-78	E156318.1	EST_HUMAN	QV0-HT0367-150200-114-g09 HT0367 Homo sapiens cDNA
3083	13010		2.22	2.0E-78		NT	Homo sapiens type IV collagen alpha 5 chain (COL4A5) gene, exon 20
	13841		1.68	2.0E-78		EST_HUMAN	EST182583 Jurkat T-cells VI Homo sapiens cDNA 5' end
	16274	26435	1.46	2.0E-78		EST_HUMAN	UI-HF-BK0-aaj-g-10-0-UI.r1 NIH_MGC_36 Homo saplens cDNA clone IMAGE:3054139 5
<u> </u>	16274	26436	1.48	2.0E-78	W 402308.1	EST HUMAN	UI-HF-BK0-aaj-g-10-0-UI.11 NIH_MGC_36 Homo sapiens cDNA clone IMAGE:3054139 5
6547	16405	26584	3.88	2.0E-78 B	F689800.1	EST_HUMAN	802188529F1 NIH_MGC_49 Homo saplens cDNA clone IMAGE:4298599 5
	16587	26761	2.73	2.0E-78	AV714177.1	EST_HUMAN	AV714177 DCB Homo sapiens cDNA clone DCBAW F09 5
<u> </u>	16789	26991	1.84	2.0E-78	AJ557509.1	EST_HUMAN	Pt2.1_16_B07.r tumor2 Homo sapiens cDNA 3'
<u></u>		26892	28.	2.0E-78	AI557509.1	EST HUMAN	Pt2.1_16_B07.r tumor2 Homo sapiens cDNA 3'
İ _	-						qi50h05.x1 NCI_CGAP_BrnZ5 Homo saplens cDNA clone IMAGE:1859961 3' similar to WP:R90.1
		28559	4.5	2.0E-78	Al197837.1	EST HUMAN	CEG6325 PROTEIN KINASE
8467	18340	28605	3.28	2.0E-78	N66951.1	EST_HUMAN	za48f12.s1 Soares fetal liver spleen 1NFLS Homo sapiens cUNA clone IMACE: 283823 3
5247	15170	24943	2.9	1.0E-78		NT	Homo sapiens GAP-like protein (LOC51306), mRNA
١.	16622		1.68	1.0E-78	U52373.1	NT	Human serine/threonine kinase MNB (mnb) mRNA, complete cds
9189	18924	25350	5.14	1.0E-78	11430460 NT	LΝ	Homo sapiens low density lipoprotein-related protein 2 (LRP2), mRNA
L	18986	25327	1.26	1.0E-78		뒫	Homo sapiens similar to lymphocyte activation-associated protein (H. sapiens) (LOC63140), mRNA
L		24274	4.48	9.0E-79	11525891 NT		Homo sapiens peptide YY (PYY), mRNA
_	14643	24431	2.48	9.0E-79	BE000837.1	EST_HUMAN	RC2-BN0074-090300-014-c12 BN0074 Home sapiens cDNA
	15255	25078	12.03	9.0E-79	AB028070.1	NT	Homo sapiens mRNA for activator of S phase (kinase, complete cds
	16733	25844	2.18	9.0E-79	5454145 NT	_ <u>F</u>	Homo sapiens ubiquitin-conjugating enzyme E2E 3 (homologous to yeast UBC4/5) (UBE2E3) mRNA
1	17120	27315	4.99	9.0E-79	J02853.1	F	Homo sapiens casein kinase II alpha subunit niRNA, complete cds
L	17120	27316	4.89	ł	J02853.1	M	Homo sapiens casein kinase II alpha subunit mRNA, complete cds
	17803	28042	1.31	9.0E-79	AF062346.1	NT	Homo sapiens zinc finger protein 216 splice variant 1 (ZNF216) mRNA, complete cdo
_	17803	28043	1.31	9.0E-79	AF062346.1	NT	Homo sapiens zinc finger protein 216 splice variant 1 (ZNF216) mRNA, complete cds
L	18293	28547	1.82	9.0E-79	AY0082	NT.	Homo sapiens TRAF6-regulated IKK activator 1 beta Uev14 mRNA, complete cds
	18616	28906	2.82	9.0E-79		Z	Homo sapiens suppressor of white apricot homolog 2 (SWAP2), mRNA
	18616	28907	2.82	9.0E-79		Ŋ	Homo sapiens suppressor of white apricot homolog 2 (SWAP2), mRNA
<u> </u>	19347	25213	1.49	9.0E-79	11417877 NT	NT	Homo sapiens gamma-glutamyttransferase 1 (GGT1), mRNA

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Top Hit Descriptor	Homo sapiens chromosome 21 segment HS21 C010	Human mRNA for KIAA0045 gene, complete cds	Human mRNA for KIAA0045 gene, complete cds	Homo saplens period (Drosophila) homolog 3 (PER3), mRNA	601472766T1 NIH_MGC_68 Homo sapiens cDNA clone IMAGE:3875657 3'	pg4e04.s1 Soares_fetal_liver_spleen_1NFLSI_S1 Homo sapiens cDNA clone IMAGE:462558 3' similar to TR:Q15408 Q15408 NEUTRAL PROTEASE LARGE SUBUNIT;	Homo sapiens chromosome 21 segment HS21C082	Homo saplens hypothetical protein FLJ10283 (FLJ10283), mRNA	601874522F1 NIH_MGC_54 Homo sapiens cDNA clone IMAGE:4101245 5'	Homo sepiens intersectin short Isoform (ITSN) mRNA, complete cds	Homo sapiens cell-line tsA201a chloride Ion current Inducer protein I(Cin) gene, complete cds	Human zinc finger protein ZNF131 mRNA, partial cds	Homo saplens MSTP016 (MST016) mRNA, complete cds	Homo saplens mRNA for KIAA0892 protein, partial cds	Homo sapiens netrin 1 (NTN1), mRNA	Homo saplens netrin 1 (NTN1), mRNA	Homo sapiens mRNA for KIAA0620 protein, partial cds	Homo sapiens mRNA for KIAA0620 protein, partial cds	601159415F2 NIH_MGC_53 Homo sapiens cDNA clone IMAGE:3511107 5'	Homo sapiens BCL2-like 2 (BCL2L2) mRNA	th18h07.x1 NCL_CGAP_Pr28 Homo sapiens cDNA clone IMAGE:2118685 3'	Homo sapiens phosphodiesterase 6A, cGMP-specific, rod, alpha (PDE6A), mRNA	Homo sapiens phosphodiesterase 8A, cGMP-specific, rod, alpha (PDE6A), mRNA	Homo sapiens mRNA for Fas-associated factor, FAF1 (Faf1 gene)	Homo sapiens hepatocellular carcinoma-associated antigen 88 (HCA88) mRNA, complete cds	Homo sapiens chloride channel CLC4 (CIC4) niRNA, complete cds	Homo saplens mRNA for Fas-associated factor, FAF1 (Faf1 gene)	Homo saplens chromosome 21 segment HS21006	Homo sapiens Rho GTPase activating protein 6 (ARHGAP6), transcript variant 4, mRNA	Homo saplens Rho GTPase activating protein 6 (ARHGAP6), transcript variant 4, mRNA	Homo sapiens hypothetical protein FLJ11006 (FLJ11006), mRNA	H4(D10S170)=putative cytoskeletal protein [hurnan, thyroid, mRNA, 3011 nt]	H4(D10S170)=putative cytoskeletal protein (fruman, thyroid, mRNA, 3011 nt)
Top Hit Database Source	LN	L	N	N.	EST_HUMAN	EST_HUMAN	LN.	NT	EST_HUMAN	LN LN	IN	TN	TN	NT	LΝ	NT	LΝ	LN LN	EST_HUMAN	NT	EST_HUMAN	IN	NT	NT	NT	Ŋ	N	NT	NT	N	LN.	NT	LN T
Top Hit Acession No.	AL163210.2	D28476.1	D28476.1	8567387 NT	BE619648.1	AA699829.1	AL163282.2	8922325 NT	BF210869.1	AF114488.1	AF232708.1	U09410.1	AF110322.1	AB020699.1	11426770 NT	11426770 NT	AB014520.1	AB014520.1	BE379926.1	4757841 NT	AI523747.1	4585863 NT	4585863 NT	AJ271408.1	AF244138.1	AF170492.1	AJ271408.1	AL163206.2	7382479 NT	7382479 NT	11427428 NT	S72869.1	S72869.1
Most Similar (Top) Hit BLAST E Value	8.0E-79	8.0E-79	8.0E-79	8.0E-79	7.0E-79	6.0E-79		4.0E-79	4.0E-79	3.0E-79	3.0E-79	3.0E-79	3.0E-79	3.0E-79	3.0E-79	3.0E-79	3.0E-79	3.0E-79	2.0E-79	2.0E-79	2.0E-79	2.0E-79	2.0E-79	2.0E-79	2.0E-79	2.0E-79	20E-79	2.0E-79	2.0E-79	2.0E-79	2.0E-79	20E-79	2.0E-79
Expression Signal	12	0.94	0.94	4.1	6.85	1.29	4.15	1.12	1.33	4.1	2.68	1.36	4.52	1.71	3.47	3.47	3.28	3.28	1.05	1.14	1.43	14.14	14.14	66.0	1.54	0.86	1.17	98'0	1.83	1.83	2.6	1.44	1.44
ORF SEQ ID NO:	23382			24889	22941		28891		24590	20091	20733		25012	25486		25508	16097	26092	20367	20685		21888	21889		52049	23542	23765		28270	26271	27029		27897
Exen SEQ ID NO:	13596	14287	14287	16097	13138	18827	<u>l</u> _		14823	10272	10885	12987	15212	15424	l	1		15960	10555		10937	11890	11990	12034	12149	13749	13976	14478	16117	16117	L	17658	17658
Probe SEQ ID NO:	3682	4391	4391	9107	3214	9040	8786	3139	4948	310	982	3060	5291	9099	5525	5525	6200	6200	618	912	1019	2101	2101	2148	2265	3838	4074	4590	6251	6251	6956	7808	7808

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	Top Hit Descriptor	RC4-BT0310-110300-015-110 BT0310 Homo sapiens cDNA	RC4-BT0310-110300-015-f10 BT0310 Homo sapiens cDNA	Homo sapiens KIAA0879 protein (KIAA0879), inRNA	Homo sapiens mRNA for KIAA0833 protein, pertial cds	Homo sapiens cadherin EGF LAG seven-pass G-type receptor 1 (CELSR1), mRNA	MR0-NN0087-260600-017-b10 NN0087 Hom; saplens cDNA	QV2-HT0540-120900-358-a05 HT0540 Homo sapiens cDNA	ai23e05.s1 Soares_testis_NHT Homo sapiens cDNA clone 1343648 3'	ai23e05.s1 Soares_testis_NHT Homo sapiens cDNA clone 1343648 3'	601581652F1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:3936061 5	Homo sapiens solute carrier family 7 (cationic amino acid transporter, y+ system), member 8 (SLC7A8), mRNA	Homo sapiens solute carrier family 7 (cationic amino acid transporter, y+ system), member 8 (SLC7A8), mRNA	Homo saplens Y chromosome spermatogenesis candidate protein (RBM) pseudogene mRNA, partial cds	Homo sapiens KIAA0724 gene product (KIAA0724), mRNA	Homo sapiens KIAA0724 gene product (KIAA0724), mRNA	Homo sapiens triple functional domain (PTPRF Interacting) (TRIO), mRNA	Homo sapiens triple functional domain (PTPRF interacting) (TRIO), mRNA	#58402.X1 NCI_CGAP_Bm23 Homo sapiens cDNA clone IMAGE:2103459 3' similar to SW:NUEM_HUMAN Q16795 NADH-UBIQUINONE OXIDOREDUC TASE 39 KD SUBUNIT PRECURSOR;	Homo sapiens NRD convertase mRNA, complete cds	Homo sapiens mRNA for KIAA1155 protein, partial cds	Homo saplens mRNA for KIAA1155 protein, partial cds	Homo sapiens malate dehydrogenase 2, NAD (initochondrial) (MDH2), mRNA	Homo sapiens mRNA for dynein heavy chain (DNAH9 gene)	Homo saplens tubby like protein 3 (TULP3), mRNA	Homo saplens G protein-coupled receptor 51 (GPR51), mRNA	Homo saplens G protein-coupled receptor 51 (GPR51), mRNA	Homo saplens chromosome 21 segment HS210101	Human cone photoreceptor cGMP-phosphodiesterase alpha' subunit gene, exon 21	Homo sapiens brefeldin A-inhibited guanine nucleotide-exchange protein 1 (BIG1), mRNA
21 1125	Top Hit Database Source	EST_HUMAN	EST_HUMAN	N _T	NT	N	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	Į.	IN	LN LN	N-	N	NT	EST_HUMAN	IN	TN	NT	NT	NT	NT	NT	NT	NT	NT	۲
	Top Hit Acession No.	BE064386.1	BE064386.1	7682357 NT	AB020640.1	11418322 NT	BF363071.1	BF087405.1	AA725848.1	AA725848.1	BE798603.1	11433924 NT	11433924 NT	U94387.1	11422647 NT	11422647 NT	6005921 NT	6005921 NT	A1422197.1	U64898.1	AB032981.1	AB032981.1	11421462 NT	AJ404468.1	11436736 NT	11526464 NT	11526464 NT	.2	U20211.1	11427366 NT
	Most Similar (Top) Hit BLAST E Value		2.0E-79	2.0E-79	2.0E-79	2.0E-79	1.0E-79	1.0E-79	9.0E-80		9.0E-80	9.0E-80	9.05-80		8.0E-80	8.0E-80	8.0E-80	8.0E-80	6.0E-80	6.0E-80		6.0E-80	6.0E-80	6.0E-80	6.0E-80	6.0E-80	6.0E-80	6.0E-80	8.0E-80	6.0E-80
	Expression Signal	4.22	4.22	2.6	4.23	1.98	2.78	2.74	3.79	3.79	1.28	11.05	11.05	0.95	2.83	2.83	1.19	1.19	2.34	2.05	1.09	1.09	4.01	2.56	3.84	3.07	3.07	1.74	1.68	2.91
	ORF SEQ ID NO:	28516	28517	24888	25341	25307		28998	22828	22829	27824	28735	28736		26512				20657	21386		23868		25712		27191	27192			28437
	Exan SEQ ID NO:	18267	18267	15098	18907	19050	19456	18704	13033	13033	17601	18464	18464	1	16343	16343		17276	10808	11528	14088	14088	15461	15610	15683	17000	17000	17091	17522	18188
	Probe SEQ ID NO:	8391	8391	8078	9161	9391	5971	8894	3107	3107	7751	8597	8597	3551	6485	6485	7409	7409	882	1624	4188	4188	5545	2025	5776	7123	7123	7214	7672	8311

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Table 4
Single Exon Probes Expressed in Heart

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Top Hit Descriptor	Homo sapiens Cy119 mRNA, complete cds	#58402.x1 NCI_CGAP_Bm23 Homo saplens cDNA clone IMAGE::2103459 3' similar to SW:NUEM_HUMAN Q16795 NADH-UBIQUINONE OXDOREDUCTASE 39 KD SUBUNIT PRECURSOR;	Homo sapiens glutathione S-transferase theta/2 (GSTT2) and glutathione S-transferase theta 1 (GSTT1) genes, complete cds	Homo sapiens CST gene for cerebroside sulfcitansferase, exon 1, 2, 3, 4, 5	Homo capiens mRNA for sodium-glucose cotransporter (SGLT2 gene)	Homo sapiens proteasome (prosome, macropain) 26S subunit, non-ATPase, 3 (PSMD3) mRNA	Homo saplens serine-threonine protein kinase (MNBH) mRNA, complete cds	Homo saplens serine-threonine protein kinase (MNBH) mRNA, complete cds	H. sapiens nox1 gene (exon 12)	Homo saplens chromosome 21 segment HS21 C083	Human ((3)mbt protein homolog mRNA, complete cds	Homo sapiens mRNA for KIAA1434 protein, partial cds	Homo saplens H3 histone family, member J (H3FJ) mRNA	Homo saplens HMT-1 mRNA for beta-1,4 mannosyltransferase, complete cds	Homo saplens HMT-1 mRNA for beta-1,4 mannosyltransferase, complete cds	Homo saplens chromosome 21 segment HS21 C068	Mus musculus keratin complex 2, gene 6g (Krt2-6g), mRNA	HSPD13155 HM3 Homo saplens cDNA clone 94000045F03	Homo sapiens chromosome 21 segment HS21,0010	PM0-GN0018-040900-002-E03 GN0018 Homo sapiens cDNA	QV4-BN0263-040600-241-g10 BN0263 Homo sapiens cDNA	co23e12.x1 Soares_NSF_F8_9W_OT_PA_P_S1 Homo sapiens cDNA clone IMAGE:1567054 3' similar to	TR:035780 035780 PIG-L.;	lyg65a08.r1 Soares infant brain 1NIB Homo sapilens cDNA clone IMAGE:38060 5'	RET4B7 subtracted retina cDNA library Homo saplens cDNA clone RET4B7	DKFZp434D1323_r1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434D1323 5'	Homo sapiens Golgi transport complex protein (90 kDe) (GTC90), mRNA	270112.r1 Sogres_testis_NHT Homo sapiens cDNA clone IMAGE:727727 5' similar to TR:G191315	Homo saplens chronosome 21 segment HS21C103	Union employe approximately 21 information mDMA	ראורווו זייינוויים אינויים ביו שנייים אינויים ביו שנייים אינויים מוויים פוויים פוויים פוויים פוויים פוויים ביו
Top Hit Database Source	R	EST_HUMAN	N	N.	N	TN	TN	NT	IN.	N	TN	N	INT	N	NT	NT	NT.	EST_HUMAN	NT .	EST_HUMAN	EST_HUMAN		EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	HOL	NAMOU I DE		IN.
Top Hit Acession No.	AF226730.1	A1422197.1	AF240786.1	AB029900.1	AJ133127.1	4508228 NT	AF108830.1	AF108830.1	X91647.1	AL163283.2	U89368.1	AB037855.1	4504292 NT	AB019038.1	AB019038.1	AL163268.2	9810293 NT	F25915.1	AL163210.2	BF085009.1	BE817465.1		A1091675.1	R35321.1	A1444821.1	AL043116.2	11421930 NT	A A DODODO 4	AL 163303.1	A CO040014	AF 23 1920. 1
Most Similar (Top) Hit BLAST E Velue	6.0E-80	6.0E-80	6.0E-80	6.0E-80	6.0E-80	. 5.0E-80	5.0E-80		5.0E-80				5.0E-80				5.0E-80		3.05-80		3.05-80			2.0E-80	2.0E-80		2.0E-80			20 10 1	
Expression Signal	50.71	1.98	1.62	4.55	2.25	1.11	1.26	1.26	4.77	2.28	0.92	3.98	1.13	1.07	1.07	1.74	1.48	7.48	4.71	1.43	4.97		2.05	6.39	1.91	6.6	1.58	96	2.00	157	7:17
ORF SEQ ID NO:	28694	20657				20319	20202				22091	22152	22518	23636		24531	8692	27441		24287						21791			0/337	79900	
Exan SEQ ID NO:	18425	10808	19562	19037	19686	10511	10745	10745	11084	11345	12192	12260	12626	13861		14752	16745	17237	10181	14498	14701		15470	11656	11713	11901	15979	}	1020		10/16
Probe SEQ ID NO:	8555	9047	9172	2871	9847	573	817	817	1172	1440	2311	2380	2784	3953	3953	4872	9989	7333	210	4610	4818		5554	1757	1816	5000	6132	25	337	i i	70/

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Top Hit Acession Top Hit Descriptor Top Hit Descriptor Source	Inn01f12.x5 NOL_CGAP_Co9 Homo sapiens cDNA clone IMAGE:1076495 3' similar to contains OFR.t1 OFR AI732858.1 EST_HUMAN repetitive element;	AF077188.1 NT Homo sapiens cullin 4A (CUL4A) mRNA, complete cds		BE386815.1 EST_HUMAN 601274305F1 NIH_MGC_20 Homo sepiens ciDNA clone IMAGE:3615433 5	L10347.1 NT Human pro-alpha1 type II collegen (COL2A1) gene exons 1-54, complete cds	Homo sapiens malate dehydrogenase 2, NAD (mitochondrial) (MDH2), nuclear gene encoding mitochondrial protein, mRNA	HUMAN	Al948731.1 [EST_HUMAN wq25c05.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2472298 3'	AF245219.1 NT Homo sapiens probable mannose binding C-type lectin DC-SIGNR mRNA, complete cds	AF24529.1 INT Homo sapiens probable mannose binding C-type lectin DC-SIGNR mRNA, complete cds	D63479.2 NT Homo sapiens mRNA for KIAA0145 protein, partial cds	11641276 NT Homo saplens similar to ret myomegalin (LOC¢4182), mRNA	11641276 NT Homo sapiens similar to rat myomegalin (LOC94182), mRNA	11417901 NT Homo saplens meringioma (disrupted in balanced translocation) 1 (MN1), mRNA		1 NT	EST_HUMAN	AI251752.1 EST_HUMAN qh90g05x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:1854298 3'		AI822115.1 EST_HUMAN ze91c08.x5 Soares_fetal_lung_NbHL19W Homo sapiens cDNA clone IMAGE:299918 3'		T_HUMAN		4501848 NT Homo sapiens ATP-binding cassette, sub-family A (ABC1), member 3 (ABCA3), mRNA		BF679022.1 EST_HUMAN 602153656F1 NIH_MGC_83 Homo saplens GiVA clone IMAGE:4294601 5'	BF679022.1 EST_HUMAN 602153666F1 NIH_MGC_83 Homo capiens cipNA clone IMAGE:4294601 5	BE288042.1 EST_HUMAN 601125505F1 NIH_MGC_B Homo sapiens cDNA clone IMAGE:3345480 5'		AB007923.1 NT Homo sapiens mRNA for KIAA0464 protein, pértial cds	08634 NT	AF252257.1 NT . Homo sapiens CRP2 binding protein mRNA, partial cds
Top Hit Acession No.						5174540						11641276	11641276	11417901									4501848	4501848							06634	
Most Similar (Top) Hit BLAST E Value	1.0E-80 P		1.0E-80[/	1.0E-80	1.0E-80 L	1.0E-80	1.0E-80 A		1.0E-80	1.0E-80	1.0E-80	1.0E-80	1.0E-80	1.0E-80	1.0E-80 A		8.0E-81 A	8.0E-81	8.0E-81	_		_	6.0E-81	6.0E-81			6.0E-81	5.0E-81		6.0E-81 ₽		4.0E-81 A
Expression Signal	3.13	96'0	0.97	5.63	6.41	1.8	2.68	2.68	1.23	1.23	1.19	7.42	7.42	1.45	1.86	1.99	2.33	2.33	4.95	3.06	4.84	4.84	1.93	1.93	1.34	1.82	1.82	3.14	3.47	3.47	2.51	0.87
ORF SEQ ID NO:		24039	24819		25635	25947	28488	28497	27417	27418	28049	28183	28184	25281	25258		28212	28213	28631	26321	23971	23872	24920	24921	27401	25269	26270	21960	26974	26975	28966	19998
Exan SEQ ID NO:	11806	14254		15187	15547	15822	16329	16329	17218	17218	17807	17935	17935	18079			17982	17982	18367	16164	14188			15153	17201	19166	19166	12057	16780	16780		10185
Probe SEQ ID NO:	1911	4358	5192	5295	5633	5916	8470	6470	7350	7350	7887	8044	8044	9443	9643	9670	8071	8071	8494	6300	4290	4290	5229	5229	7325	8258	6296	2170	6901	6901	8865	214

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	Top Hit Descriptor	EST372729 MAGE resequences, MAGF Horrio sapiens cDNA	2k45h09.r1 Soares, pregnant, uterus, NbHPU Homo saptens cDNA clone IMAGE:485825 5' similar to PIR:S52437 S52437 CDP-diacyglycerol synthase - fruit fly;	1245c04.y1 NCI_CGAP_Brn52 Homo sepiens cDNA clone IMAGE:2291526 5'	Human aconitate hydratase (ACO2) gene, exon 3	Homo sapiens polymerase (DNA directed), gamma (POLG), mRNA	Homo sapiens polymerase (DNA directed), gamma (POLG), mRNA	Homo sapiens arm-repeat protein NPRAP/neurojungin (CTNND2) mRNA, partial cds	Homo sapiens arm-repeat protein NPRAP/neurojungin (CTNND2) mRNA, partial cds	602137864F1 NIH_MGC_83 Homo sapiens cDNA clone IMAGE:4274635 5'	Homo sapiens polymerase (DNA directed), gaimma (POLG), mRNA	601645051F1 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:3930228 5	601645051F1 NIH_MGC_56 Homo sapiens dDNA clone IMAGE:3930228 5'	601343180F1 NIH_MGC_53 Home sapiens cDNA clone IMAGE:3685483 5	601577339F1 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:3838280 5'	601577339F1 NIH_MGC_9 Homo saplens cDNA clone IMAGE:3838280 5'	CM3-NN0059-140400-147-a12 NN0059 Homo sapiens cDNA	MR0-CT0006-250599-019 CT0006 Homo sapiens cDNA	MR0-CT0008-250599-019 CT0006 Hamo saplens cDNA	EST372729 MAGE resequences, MAGF Homb saplens cDNA	601867714F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:4110459 5'	Homo saplens phorbolin (similar to apolipoprotein B mRNA editing protein) (DJ742C19.2), mRNA	Homo saplens HSPC288 mRNA, partial cds	Homo sapiens HSPC288 mRNA, partial cds	Human CRFB4 gene, partial cds	Human CRFB4 gene, partial cds	Human CRFB4 gene, partial cds	Homo sapiens mRNA for KIAA1327 protein, pairtial cds	Homo sapiens glutathione peroxidase 5 (epididymal androgen-related protein) (GPX5), transcript variant 2,	IMRNA	Homo sapiens hypothetical protein FLJ20461 (FLJ20461), mRNA	601458531F1 NIH_MGC_66 Homo saplens cDNA clone IMAGE:3862086 5	AU144050 HEMBA1 Homo sapiens cDNA clone HEMBA1000752 3'	Inf69e11.s.1 NCI CGAP Co3 Home septems cDNA clone IMAGE 925196.3
	Top Hit Database Source	EST_HUMAN	EST HUMAN	EST_HUMAN	L	L	NT	TN	INT	EST_HUMAN	LN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	NT	INT	IN	INT	IN	IN	NT		NT	NT	EST_HUMAN	EST HUMAN	FST HIMAN
	Top Hit Acesslon No.	AW960658.1	AA040370.1	BE047996.1	U87928.1	11432966 NT	11432966 NT	U52351.1	U52351.1	BF674641.1	11432968 NT	BE958278.1	BE958278.1	BE564367.1	BE744545.1	BE744545.1	AW897650.1	AW844986.1	AW844986.1	AW960658.1	BF204253.1	11418138 NT	AF161406.1	AF161406.1	U08988.1	U08988.1	U08988.1	AB037748.1		6715601 NT	8923432 NT	BF035327.1	AU144050.1	AA515512 1
	Most Similar (Top) Hit BLAST E	1.0E-81	1.0E-81	_	1.0E-81	1.0E-81	1.0E-81	1.0E-81			1.0E-81			1.0E-81	1.0E-81	1.0E-81	1.0E-81	1.0E-81	1.0E-81		1.0E-81	1.0E-81	8.0E-82	8.0E-82	8.0E-82	8.0E-82	8.0E-82	8.0E-82		8.0E-82		7.0E-82		5 OF-82
	Expression Signal	2	3.07	7.85	9.03	4.01	4.01	3.54	3.54	3:15	6.4	2.62	2.62	4.81	2.93	2.83	1.46	1.98	1.98	2.42	1.98	3.62	1.87	1.28	1.66	2.17	1.11	1.12		1.24	0.81	1.27	1.82	0.81
	ORF SEQ ID NO:	23304	24091		29103	24982	24983	25449	28450		26606		27703	27799	27809		28091	28568			28915	25325	19789	19789	20044	20566	20645	21244		21400	23825		22495	
	SEQ ID NO:	13516	14308		15083	15206	15208	15388	15388	15645	16425	17482	17482	17576	17669	17669	17850	18312	18312	13516	18624	18981	8666	8668		10726	10795	11380		11540	14050	11339	12601	13937
	Probe SEQ ID NO:	3602	4414	4546	5203	5284	5284	5468	8468	2232	2929	7631	7631	7726	7819	7819	8000	8438	8438	8694	8810	9278	12	101	263	797	889	1475		1636	4150	1434	2739	4034

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Top Hit Descriptor	Homo sapiens alpha-tubulin isoform 1 mRNA, complete cds	wp75e09.x1 NCI_CGAP_Bm25 Homo sapiens cDNA clone IMAGE:2467624 3' similar to TR:075276 075276 PKD1;	Homo sapiens presentiin-1 gene, exons 1 and 2	Homo sapiens amylaid beta (A4) precursor protein (protease nexin-II, Alzheimer disease) (APP), mRNA	RC2-BN0120-010400-013-f02 BN0120 Homo sapiens cDNA	Homo saplens transforming growth factor beta-activated kinase-binding protein 1 (TAB1), mRNA	Homo sapiens amykad beta (A4) precursor prolein (protease nexin-ll, Alzhelmer disease) (APP), mRNA	al23e05.s1 Soares_testis_NHT Homo sapiens cDNA clone 1343648 3'	RC6-PT0001-190100-021-B02 PT0001 Homo sapiens cDNA	Homo sapiens chromosome 21 segment HS21,0085	RC1-BN0005-260700-018-g04 BN0005 Homo sapiens cDNA	Homo saplens adenylate cyclase activating polypeptide 1 (pltultary) receptor type I (ADCYAP1R1) mRNA	Homo sapiens neurotrophic tyrosine kinase, receptor, type 2 (NTRK2) mRNA	Zn93b04.r1 Stratagene lung carcinoma 937218 Homo saplens cDNA clone IMAGE:565711 5' similar to SW:PAGT_BOVIN Q07537 POLYPEPTIDE N-ACETYLGALACTOSAMINYLTRANSFERASE;	Homo sapiens ankyrin-like with transmembrane domains 1 (ANKTM1), mRNA	Homo sapiens mRNA for KIAA 1077 protein, partial cds	Homo saplens mRNA for KIAA1077 protein, partial cds	Homo sapiens mRNA for KIAA0899 protein, partial cds	Homo sapiens mRNA for KÍAA0999 protein, partial cds	DKFZp434M117_r1 434 (synonym: htes3) Horrio sapiens cDNA clone DKFZp434M117 5'	Homo sapiens DNA for amyloid precursor protein, complete cds	Homo sapiens glutamate receptor, lonotropic, kainate 1 (GRIK1) mRNA	Hamo sapiens mRNA for KIAA1096 protein, partial cds	Homo sapiens mRNA for KIAA1096 protein, partial cds	Homo sapiens wbscr1 (WBSCR1) and wbscr5 (WBSCR5) genes, complete cds, elternatively spliced and replication factor C subunit 2 (RFC2) cene, complete cds	Homo sepiens tumor necrosis factor receptor superfamily, member 5 (TNFRSF5) mRNA	Homo sapiens tumor necrosis factor receptor superfamily, member 5 (TNFRSF5) mRNA
Top Hit Database Source	NT	EST_HUMAN	LN.	NT	EST_HUMAN	NT	TN	EST_HUMAN	EST_HUMAN	LΝ	EST_HUMAN	LΝ	NT	EST_HUMAN	۲	F	LN	NT	NT	EST_HUMAN	NT	NT	TN	NT	FZ	NT NT	L
Top Hit Acession No.	AF081484.1	A1937300.1	AF029701.2	4502166 NT	BE005705.1	5174702 NT	4502166 NT	AA725848.1	AW875073.1	AL163285.2	BE813232.1	4501922 NT	5453811 NT	AA135979.1	11425208 NT	AB029000.1	AB029000.1	AB023216.1	AB023216.1	AL046390.1	D87675.1	4504116 NT	AB029019.1	AB029019.1	A F045555 1	4607580 NT	4507580 NT
Most Similar (Top) Hit BLAST E Value	4.0E-82		4.0E-82	3.0E-82	3.0E-82	3.0E-82	3.0E-82	3.0E-82		3.0E-82	3.05-82	3.0E-82	3.0E-82	3.0E-82	3.0E-82	3.0E-82		2.05-82		2.0E-82	2.0E-82			2.0E-82		2.0E-82	2.0E-82
Expression Signal	6.51	6.47	6.19	13.75	2.19	4.44	3.22	13.76	5.47	2.03	1.82	6.0	2.06	0.91	2.84	3.79	3.79	1.92	1.82	1.75	1.14	1.14	86.0	0.98	3.18	1.66	1.68
ORF SEQ ID NO:	21412	28054		20061	20446	20539	20629		21098			21744		24501	26807	27728	27729		20327	21428	23466	23815	24143	24144	24441		24654
Exan SEQ ID NO:	11551	18761	19130	10242	10620		10779		11240	11355	11755	11855	13158	14718	16617	17504	17504		10520	11562	13684	14040	14352	14352	14653		1 1
Probe SEQ ID NO:	1647	8954	9519	278	289	770	852	1045	1333	1450	1859	1961	3234	4836	6733	7654	7654	285	582	1660	3772	4140	4458	4458	4768	6013	5013

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Гор Hit Descriptor	Homo saplens mRNA for KIAA0727 protein, partial cds	Homo sapiens FAM4A1 splice variant a (FAM4A1) mRNA, complete cds	Homo sapiens slit (Drosophila) homolog 3 (SLIT3), mRNA	Human endogenous retrovirus-K, LTR U5 and gag gene	Human endogenous retrovirus-K, LTR U5 and gag gene	Homo saplens CAGF9 mRNA, partial cds	Homo saplens CAGF9 mRNA, partial cds	zb31d10.s1 Scares_parathyroid_turnor_NbHPA Homo sapiens cDNA clone IMAGE:305203 3'	201g09.r1 Scares fetal_liver_spleen_1NFLS_iS1 Homo saplens cDNA clone IMAGE:429569 5	Homo sapiens SRY (sex determining region Y)-box 10 (SOX10), mRNA	Homo saplens melanoma differentiation associated protein-5 (MDA5), mRNA	601510859F1 NIH_MGC_71 Homo saplens clONA clane IMAGE:3912207 5'	RC4-BT0310-110300-015-110 BT0310 Homo sapiens cDNA	Homo sapiens mRNA for KIAA0538 protein, partial cds	UI-H-BW1-aca-f-03-0-UI.s1 NCI_CGAP_Sub? Homo sapiens cDNA clone IMAGE:3084053 3	Homo sapiens chromosome 21 segment HS210009	Homo saplens chramosome 21 segment HS21C046	802150403F1 NIH_MGC_81 Homo sapiens d2NA clane IMAGE:4291561 5	601273346F1 NIH_MGC_20 Homo saplens clONA clone IMAGE:3814362 5	za48f12.s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:295823 3'	no12h01.s1 NCL_CGAP_Phe1 Homo sepiens cDNA clone IMAGE:1100497 3' sImilar to contains Alu repetitive element;	7p37a07.x1 NCI_CGAP_Pr28 Homo sapiens cDNA clone IMAGE:3647893 3' similar to TR:Q9Y316 Q9Y316	DJ207H1.1;	Human platelet Glycoprotein IIb (GPIIb) gene, exons 2-29	hf31h03.x1 Soares_NFL_T_GBC_S1 Homo scipiens oDNA clone IMAGE:2933525 3' similar to	OWLECTOR 44100 032 66 ST034 Home inclined in Control	CATAC CONTROL COLOR CATACON COLOR CATACON COLOR CATACON COLOR CATACON COLOR CATACON COLOR CATACON CATA	Z59c05,s1 Sogres fetal liver spieen 1NFLS S1 Homo saplens cDNA clone IMAGE:435080 31	Homo sapiens hypothetical protein FLJ10379 (FLJ10379), mRNA	Homo saplens VAMP (vesicle-associated membrane protein)-associated protein A (33kD) (VAPA) mRNA,	and translated products	Homo sapiens hyperion gene, exons 1-50	Homo sapiens met proto-oncogene (hepatocyte growth factor receptor) (MET), mRNA
Top Hit Database Source	NT	N	NT	TN	NT	NT	NT	EST_HUMAN	EST_HUMAN	TN	NT	EST_HUMAN	EST_HUMAN	TN	EST_HUMAN	TN	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN		EST_HUMAN	NT	MAN ILL FOR	COT TOTAL	אואוחם בו כש	EST HUMAN	ΙN		N	Ł	NT
Top Hit Acession No.	AB018270.1	AF234882.1	11321570 NT	Y08032.1	Y08032.1	U80736.1	U80736.1	N94950.1	AA011278.1	11418097 NT	11545921 NT	BE885106.1	BE064386.1	AB011110.2	BF515938.1	AL163209.2	AL163246.2	BF672220.1	BE383973.1	N66951.1	AA584655.1		BF221813.1	M33320.1	A 18/ E79/00 4	A14/040404.4	AW 6 10403. I	AA701457.1	11430241 NT		4507866 NT	AJ010770.1	11422024 NT
Most Similar (Top) Hit BLAST E Value	$\overline{}$		2.0E-82	2.0E-82	2.0E-82	2.0E-82	2.0E-82	2.0E-82		2.0E-82	1.0E-82	1.0E-82	1.0E-82	1.0E-82	1.0E-82	1.0E-82			I	8.0E-83	7.0E-83	_		6.0E-83		0.00-00	0.0E-00	6.0E-83	6.0E-83		6.0E-83		6.0E-83
Expression Signal	2.37	4.96	2.23	1.2	1.2	6.98	6.98	2.23	2.57	1.44	1.45	6.0	1.91	0.83	1.38	2.57	1.76	4.7	1.88	1.96	1.64		5.49	1.34	4	2 6	BO:	0.94	0.95		1.72	2.13	1.79
ORF SEQ ID NO:	H			27912	27913	28766	28767				20321		21020	21021		28264				21422				20169					23231				26452
Exon SEQ ID NO:	LI	_	16713	17871	17871	18493	18493	18868	19205	19395	10515	11100	11171	11172	17732	18016			11297	12647	12764	l	ı	10343	14044	100,	7187	- (13431		- 1	- 1	16291
Probe SEQ ID NO:	5358	5748	6834	7821	7821	8628	8628	2608	9632	6923	222	1190	1264	1265	7882	8128	8373	7057	1392	1656	2836		4710	397	1717	/4/	1000	3331	3515		5236	5669	8430

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Probe SEQ ID	Exen SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
7579	17430	27644	6.77	6.0E-83	4505314 NT	٦	Homo sapiens mycmesin (M-protein) 2 (165kD) (MYOM2), mRNA
8817	18630		6.32	6.0E-83	6.0E-83 AA486105.1	EST_HUMAN	ab14e10.s1 Stratagene lung (#937210) Homo sapiens cDNA clone IMAGE:840810 3' similar to contains THR.t2 THR repetitive element;
							Homo saplens glutathione S-transferase theta 2 (GSTT2) and glutathione S-transferase theta 1 (GSTT1)
9050	18834		3.11	6.0E-83	.1	NT	genes, complete cds
931	10856		4.14	5.0E-83	5.0E-83 U17883.1	NT	Human succinate dehydrogenase Iron-protein (subunit (sdhB) gene, exon 5
2004	12649		2.1	5.0E-83	5.0E-83 AF006305.1	IN	Homo sapiens 28S proteasome regulatory subunit (SUG2) mRNA, complete cds
3586	13500	23289	0.92	5.0E-83 AL	AL133207.2	TN	Novel human gene mapping to chomosome X
5011	14885	24651	10.99	5.0E-83	TN 81013 NT	LN	Homo sapiens catalase (CAT) mRNA
5011	14885	24652	10.99	5.0E-83	4557013 NT	Į	Homo sapiens catalase (CAT) mRNA
5094	14864	24739	0.86	5.0E-83 AF	083827.1	N	Homo sapiens dihydropyridine receptor alpha 2 subunit (CACNA2D1) gene, exon 11
3	100	02000			7 000700	<u> </u>	Homo saplens mannosidase, beta A, Iysosomai (MANBA) gene, and ubiquitin-conjugating enzyme E2D 3
624					4.0E-83 AF224669.1	Z	(DEZZLA) genes, compare cas
3469		23190	0		4.0E-83 BE888078.1	EST_HUMAN	601511580F1 NIH_MGC_71 Homo sapiens ciona ciona IMAGE:3913195 5
981	10904		4.5		3.0E-83 AA368311.1	EST_HUMAN	EST79542 Placenta i Homo sapiens cDNA similar to similar to endogenous retrovirus ERV9
2750	12612		1.82	3.0E-83	AA632654.1	EST HUMAN	np87c07.s1 NCI_CGAP_Thy1 Homo sapiens cDNA clone IMAGE:1133292 similar to contains THR.t2 THR repetitive element;
1759	11658	21529	0,	2.0E-83 AA	AA993492.1	EST HUMAN	ot64g05.s1 Soares, testis, NHT Homo sapiens cDNA clone IMAGE:1621592 3' similar to TR:Q92614 Q92814 MYELOBLAST KIAA0218.
1750					2 OE: 83 A A COS 4 2 4	ENT LIMAN	ot84g05.s1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1621592 3' similar to TR:Q92614 Doos44 MYELORI AST KIAAกวาล
1883			ľ		2.0E-83 N66951.1	EST HUMAN	za48112.s1 Spares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:295823 3'
2821	12750			2.0E-83 BE	BE828694.1	EST HUMAN	RC6-ET0046-280600-013-H12 ET0046 Homo saplens cDNA
3231	13155		1.82	2.0E-83	L	N	Homo sapiens sal (Drosophila)-like 1 (SALL1), mRNA
3708	13621		1	2.0E-83 AL	163202.2	LN	Homo sapiens chromosome 21 segment HS21C002
4241	14140	23915	4.47	2.0E-83 AF	AF202879.1	TN	Homo saplens hematopoietic progenitor cell antigen CD34 precursor (CD34) mRNA, partial cds
4553	14448	24230	8.13	2.0E-83	7706398 NT	۲	Homo sapiens ankyrin repeat-containing protein ASB-2 (LOC51676), mRNA
4553	14448	24231	8.13	2.0E-83	TN 868997	TN	Homo sapiens ankyrin repeat-containing protein ASB-2 (LOC51676), mRNA
5359	15279	25109	23.35	2.0E-83	11024711 NT	IN	Homo saplens myosin, heavy polypeptide 4, skeletal muscle (MYH4), mRNA
6369	15279	25110	23.35	2.0E-83	11024711 NT	TN	Homo sapiens myosin, heavy polypeptide 4, skeletal muscle (MYH4), mRNA
6398	16258				2.0E-83 AF129533.1	NT	Homo sapiens F-box protein Fbl3b (FBL3B) mRNA, partial cds
6829					2.0E-83 U66707.1	N	Rattus norvegicus densin-180 mRNA, complete cds
6839				ŀ	2.0E-83 AF011920.1	NT	Homo sapiens protein kinase CK2 catalytic subunit alpha gene, exon 1
6839	16718	26912	2.56		AF011920.1	보	Homo sapiens protein kinase CK2 catalytic subunit alpha gene, exon 1

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	7	7	Ť	Т	7	Т	7				T	T	T	T		۳	۳	T	Т	Τ			П		\Box	Т	Т	Τ"
Top Hit Descriptor	Human neural cell adhesion molecule (N-CAM) secreted Isoform mRNA, 3' end	Human neural cell adhesion molecule (N-CAM) secreted isoform mRNA, 3' end	AU117659 HEMBA1 Homo saplens cDNA clone HEMBA1001910 5'	Homo sapiens KIAA0985 protein (KIAA0985); mRNA	DKFZp547J135_r1 547 (synonym: hfbr1) Horno sapiens cDNA clone DKFZp547J135 5	DKFZp547J135_r1 547 (synonym: hfbr1) Horno sapiens cDNA clone DKFZp547J135 5	Homo sapiens gene for AF-6, complete cds	Homo saplens hydroxyacyt-Coenzyme A detrytrogenase/3-ketoacyt-Coenzyme A thiolase/enoyt-Coenzyme A hydratase (trifunctional protein), beta subunit (HADHB) mRNA	Homo saplens hydroxyacyl-Coenzyme A dehydrogenase/3-katoacyl-Coenzyme A thiolase/enoyl-Coenzyme A hydratase (trifunctional protein), beta subunit (HADHB) mRNA	601507375F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3908754 6'	Homo sapiens cell recognition molecule Caspr2 (KIAA0868), mRNA	Rattus norvegicus brain specific cortactin-binding protein CBP90 mRNA, partial cds	H.saplens gene for mitochondrial dodecenoyi-CoA delta-isomerase, exon 3	Homo sapiens amyloid beta (A4) precursor protein (protease nexin-li, Alzheimer disease) (APP), mRNA	owgbob.x/ Soares_testis_NHT Homo sapien's cONA clone IMAGE:1845431 3' similar to gb:M64241 QM PROTEIN (HUMAN);	601676023F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3958853 5'	RC2-FN0119-200600-011-g05 FN0119 Homo sapiens cDNA	RC2-FN0119-200600-011-g05 FN0119 Home saplens cDNA	ae86a03.s1 Stratagene schizo brain S11 Homo sapiens cDNA clone IMAGE:971020 3	DKFZp434H0322_r1 434 (synonym: htes3) Hamo sapiens cDNA clone DKFZp434H0322 5	al47g03.s1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA done IMAGE:1460500 3' similar to gb:M14338 VITAMIN K-DEPENDENT PROTEIN S PRECURSOR (HUMAN);	PM0-LT0019-190600-004-F02 LT0019 Hama saplens cDNA	PM4-FT0054-160600-004-e10 FT0054 Homo sapiens cDNA	ILO-BT0168-091199-139-e06 BT0168 Homo (appiens cDNA	EST96094 Testis I Homo sapiens cDNA 5' end	Homo sapiens chromosome 3 subtelomeric region	Homo sepiens regulatory factor X, 3 (influences HLA class II expression) (RFX3), mRNA	Homo sapiens mRNA for KIAA1314 protein, partial cds
Top Hit Database Source	NT	NT	EST_HUMAN	NT	EST_HUMAN	EST_HUMAN	NT	NT	N N	EST HUMAN	NT	NT	NT	FN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	LN.	N
Top Hit Acesslan No.	M22094.1		AU117659.1	11438448 NT	AL134452.1	AL134452.1	AB011399.1	4504328 NT	4504326 NT	BE883690.1	32349	AF053768.1	225822.1	4502188 NT	A1027614.1	BE901209.1	BE838864.1	BE838864.1	AA776574.1	AL042863.2	AA897339.1	BE810371.1	BE770199.1	AW369812.1	AA382811.1	AF109718.1	11428740 NT	AB037735.1
Most Similar (Top) Hit BLAST E Value	20E-83	2.0E-83	2.0E-83	2.0E-83	2.0E-83	2.0E-83	2.0E-83	1.0E-83	1.0E-83	-	1.0E-83	1.0E-83	1.0E-83	1.0E-83		7.0E-84	6.0E-84		6.0E-84	6.0E-84	6.0E-84	6.0E-84	8.0E-84		5.0E-84		5.0E-84	4.0E-84
Expression Signal	3.32	3.32	1.21	3.7	1.82	1.82	3.48	16.57	16.57	1.8	0.84	3.55	2.22	1.36	1.75	2.82	4.21	4.21	4.78	6.24	1.7	3.17	1.9	<u>4</u>	1.06	1.01	2.76	1.08
ORF SEQ ID NO:	27845	27846			28422	28423		21162	<u> </u>		L			24461							25180		L		20458	Ì_	28926	Ц
Exen SEQ ID NO:	17817	17817	17678	18105	18177	18177	19226	11295	J		1	L	L	14874		1_	11180	11180	ı	15085	15307		L	<u> </u>	L	12908	18643	11260
Probe SEQ ID NO:	79/7	79/7	7826	8223	8228	8238	29867	1390	1390	2620	3148	3783	4161	4789	6022	3727	1273	1273	2348	9029	8388	8415	8208	8821	697	2981	8830	1354

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				_		_				_				_		_				*****				g				٠	Martin - Pro	g. 4.	er -
Top Hit Descriptor		wa76c04.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMACE:2302086 3' similar to SW:NRDC_HUMAN O43847 NARDILYSIN PRECURSOR;	Homo saplens myosin light chain kinase Isoform 2 (MLCK) mRNA, complete cds	Human 2,4-dienoyl-CoA reductase gene, exons 3 and 4	Homo sapiens protein tyrosine phosphatase, receptor type, G (PTPRG), mRNA	Homo sapiens protein tyrosine phosphatase, receptor type, G (PTPRG), mRNA	Homo saplens histone deacetylase 3 (HDAC3) gene, complete cds	Homo sapiens KIAA0783 gene product (KIAA0783), mRNA	Homo sapiens mRNA for KIAA1130 protein, partial cds	Homo sapiens Bach1 protein homolog mRNA, partial cds	Homo saplens chondroitin sulfate proteoglycan 2 (versican) (CSPG2) mRNA	Homo sapiens pericentriolar material 1 (PCM1) mRNA	Novel human mRNA containing Zinc finger C2H2 type domains	Homo sapiens DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes,	complete cds)	Homo sapiens X-linked juvenile retinoschisis precursor protein (XLRS1) mRNA, complete cds	wu20d05.x1 Soares_Dieckgraefe_colon_NHCD Homo sapiens cDNA clone IMAGE:2520585 3' similar to ab:L05083 60S RIBOSOMAL PROTEIN L18A (HUMAN):	CM1-BT0795-190600-272-b08 BT0795 Homolsaplens cDNA	CM1-BT0785-190600-272-b08 BT0795 Homo saplens cDNA	Homo sapiens myelin transcription factor 1-like (MYT1-l) mRNA, complete cds	H.saplens DNA for endogenous retroviral like element	601887664F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:4121727 5'	601887664F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:4121727 5'	qm87c09.x1 NCI_CGAP_Lu5 Homo saplens cjNA clone IMAGE:1895728 3'	nae30a02.x1 Lupski_sympathetic_trunk Homo sapiens cDNA clone IMAGE:4090251 3' similar to TR:Q9UGS3 Q9UGS3 DJ756G23.1;	nae30a02.x1 Lupski_sympathetic_trunk Homo sapiens cDNA clone IMAGE:4090251 3' similar to	TK:Q8UGS3 Q9UGS3 DJ786G23.1;	Homo sapiens intersectin short isoform (ITSN) mRNA, complete cds	Homo sapiens tyrosine 3-monooxygenase/trypt/phan 5-monooxygenase activation protein, zeta polypeptide (YWHAZ) mRNA	Homo sapiens complement component 5 (C5), mRNA	вт85b11.s1 Stratagene schizo brain S11 Homo saplens cDNA clone IMAGE:1629885 3'
Top Hit Datebase	Source	EST_HUMAN		Ę	F	NT	NT	NT	TN	NT	N F	Z	N F		NT	NT	EST HUMAN	EST HUMAN	EST HUMAN	NT	Ę	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN		EST_HUMAN	NT	NT	NT	EST_HUMAN
Top Hit Acession	ġ	AI685321.1	AF069601.2	U94982.1	11386168 NT	11386168 NT	AF059650.1	11421328 NT	AB032956.1	AF026200.1	4758081 NT	5453855 NT	AL096880.1		AB026898.1	AF014459.1	A1983801.1	BE695397.1	3E695397.1	2.0E-84 AF036943.1	X89211.1	2.0E-84 BF308518.1	BF308518.1	AI298674.1	2.0E-84 BF448000.1		F448000.1	1.0E-84 AF114488.1	4507952 NT	11427631 NT	AA984379.1
Most Similar (Top) Hit	Value	4.0E-84	4.0E-84	4.0E-84	4.0E-84	4.0E-84	4.0E-84	4.0E-84	4.0E-84	3.0E-84	3.0E-84	3.0E-84	3.0E-84		3.0E-84 A	3.0E-84 A	3.0E-84		2.0E-84	2.0E-84	2.0E-84)	2.0E-84	2.0E-84	2.0E-84	2.0E-84	1	2.0E-84 B	1.0E-84	1.0E-84	1.0E-84	1.0E-84 A
Expression	Buggo	4.03	1.76	1.27	1.31	1.31	2.35	12.15	6.56	1.36	0.89	1.15	3.03		1.18	4.11	7.8	5.89	68.8	9.55	0.93	1.11	1.11	1.67	1.89		1.89	1.63	5.2	0.99	1.92
ORF SEQ	<u> </u>	21151	24529	24701				28543	28413	20094	20891	21690	21745		23251			21845			22659	24364	24365		25334			20080	20290		21026
Exon SEQ ID	ö	11294	14749	14930	15336	15336	15697	16366	18169	10275	11051	11812	11856		13458	13602	18133	11948	11948	12840	12859	14568	14568	16575	19003		18003	10271	10478	10636	11178
_ <u> </u>	Ö	1389	4869	2060	5416	5418	5791	6507	8230	313	1137	. 1917	1962		3542	3689	8253	2058	2058	2913	2932	4682	4682	6695	8308		8308	309	537	703	1271

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Top Hit Descriptor		601308008F1 NIH_MGC_44 Homo sapiens dJNA clone IMAGE:3626257 5	Homo sapiens pericentriolar material 1 (PCM1), mRNA	nw12e08.s1 NCI_CGAP_SS1 Homo sapiens cDNA clone IMAGE:1239108 3'	Homo sapians 959 kb contig between AML1 and CBR1 on chromosome 21q22; segment 1/3	DKFZp434N0323_r1 434 (synonym: htes3) Homo saplens cDNA clone DKFZp434N0323 5'	DKFZp434N0323_r1 434 (synonym: htes3) Homo saplens cDNA clone DKFZp434N0323 5'	Homo sapiens 959 kb contig between AML1 and CBR1 on chromosome 21q22; segment 1/3	uterine water channel≕28 kda erythrocyte integral membrane protein homdog (human, uterus, mRNA, 1340 nti	Novel human gene mapping to chomosome 13	Novel human gene mapping to chomosome 13	Novel human gene mapping to chomosome 13	Homo sapiens polymerase (DNA directed), alpha (POLA), mRNA	Homo sapiens NGFI-A binding protein 1 (ERG1 binding protein 1) (NAB1), mRNA	Homo sapiens nuclear transport factor 2 (placental protein 15) (PP15) mRNA	Homo sapiens ubiquitin specific protease 13 (teopeptidase T-3) (USP13) mRNA	Homo sapiens ublquitin specific protease 13 (isopeptidase T-3) (USP13) mRNA	Homo saplens purinergic receptor P2X-like 1, crphan receptor (P2RXL1), mRNA	Homo sapiens aconitase 2, mitochondrial (ACO2), mRNA	Homo sapiens chromosome 21 segment HS21,0009	Homo sapiens nuclear protein Skip mRNA, complete cds	Homo sapiens nuclear protein Skip mRNA, complete cds	Human plasminogen gene, exon 7	Human plasminogen gene, exon 7	Homo sapiens DKFZp434P211 protein (DKFZP434P211), mRNA	Homo sapiens chromosome 21 segment HS21 C080	Homo sapiens heat shock transcription factor 2 binding protein (HSF2BP), mRNA	Homo saplens chromosome 21 segment HS21C068	Homo saplens DKFZp434P211 protein (DKFZP434P211), mRNA	Homo sapiens ribosomal protein L27 mRNA, complete cds	Homo saplens MSTP030 mRNA, complete cds	Homo sapiens DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 10 (RNA helicase) (DDX10), mRNA
	Source	EST_HUMAN	IN	EST_HUMAN	N-	EST_HUMAN	EST_HUMAN	N L	Ł	Ŋ	N	N-	N	N FZ	INT	F	N _T	TN	INT	INT	INT	NT	TN	TN	NT	Z	۲	N	Į.	N	TN	NT
Top Hit Acession No.		BE392137.1	11427197 NT	AA720851.1	AJ229041.1	AL043314.2	AL043314.2	AJ229041.1	S73482.1	AL049784.1	AL049784.1	AL049784.1	8393994 NT	11430848 NT	5031984 NT	4507848 NT	4507848 NT	11417812 NT	11418185 NT	AL163209.2	U51432.1	U51432.1	M33282.1	M33282.1	7657020 NT	AL163280.2	5901979 NT	AL163268.2	7657020 NT	L05094.1	AF113210.1	11438573 NT
Most Similar (Top) Hit BLAST E	Value	1.0E-84	1.0E-84	1.0E-84	1.0E-84	1.0E-84		1.0E-84	1.0E-84	1.0E-84	1.0E-84	1.0E-84	1.0E-84	1.0E-84	1.0E-84	1.0E-84	1.0E-84	1.0E-84	1.0E-84	9.0E-85	9.0E-85	9.0E-85	9.0E-85	9.0E-85	9.0E-85	9.0E-85	8.0E-85	9.0E-85	9.0E-85	7.0E-85	7.0E-85	6.0E-85
Expression Signal	•	3.13	1.08	2.14	69.9	3.82	3.82	4.29	1.52	1.49	1.49	1.96	3.12	1.8	2.45	1.85	1.85	2.98	9.8	1.17	2.3	2.3	1.07	1.07	4.93	0.94	1.14	1.01	1.27	3.24	5.81	2.56
ORF SEQ ID NO:		21780	21964		23998	24275	24276	83888	25773	26143	26144	26236		26490		24884	24885		25330		20816		21322	21323		23832	24449	24491	21417	20877		28819
Exan SEQ ID	ö	11900	12063	13605	14216	14489	14489	14216	15665	16007	16007	16086	16276	16324	17365	15091	15091	18925	18998	10875	10974	10974	11465	11465	11554	14058	14862	14707	11554	11035	18718	18535
Probe SEQ ID	Ö	2008	2176	3691	4319	4601	4601	4823	5757	6113	6113	6220	6414	6483	7495	7639	7639	9190	9298	951	1057	1057	1560	1560	1651	4158	4778	4825	9819	1120	8910	8718

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Top Hit Descriptor	Homo sapiens chromosame 21 segment HS2;1C084	wi67h08.x1 NCI_CGAP_Kid12 Homo sapiens cDNA clone IMAGE:2398431 3' similar to contains element MSR1 repetitive element:	601591416F1 NIH MGC 7 Homo saplens cDNA done IMAGE 3945818 5'	601462817F1 NIH_MGC_67 Homo sapiens cDNA clane IMAGE:3866021 5	601462817F1 NIH_MGC_67 Homo saplens cDNA clone IMAGE:3866021 5'	601109738F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3350553 5	245f03.s1 Scares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:453245 3'	z/45f03.s1 Scares_fetal_liver_spleen_1NFLS_S1 Homo saplens cDNA clone IMAGE:453245 3'	601897003F1 NIH_MGC_19 Homo septens cIDNA clane IMAGE:4126440 5'	601897003F1 NIH_MGC_19 Homo saplens cDNA clane IMAGE:4126440 5	Homo saplens calcineurin binding protein 1 (K/AA0330), mRNA	Homo saplens calcineurin binding protein 1 (KJAA0330), mRNA	601120778F1 NIH_MGC_20 Homo sapiens dJNA clone IMAGE:2987690 5'	Homo sapiens KIAA0680 gene product (KIAA0680), mRNA	aj88f08.s1 Scares_parathyrold_tumor_NbHPA Homo saplens cDNA clone IMAGE:1403559 3'	aj88f08.s1 Scares_parathyroid_tumor_NbHPA Homo sapiens cDNA clone IMAGE:1403559 3'	Homo sapiens Text (human T-cell leukernla virus type I) binding protein 1 (TAX1BP1), mRNA	Homo sapiens galactocerebrosidase (GALC) gene, exon 15	Homo saplens RAN binding protein 7 (RANBP7), mRNA	Homo sapiens DiGeorge syndrome critical region gene 6 (DGCR6), mRNA	Homo sapiens similar to transcription factor CA150 (H. sapiens) (LOC63170), mRNA	Homo saplens similar to transcription factor CA 150 (H. saplens) (LOC63170), mRNA	Homo sapiens oxoglutarate dehydrogenase (lippamide) (OGDH) mRNA	601072594F1 NIH_MGC_12 Homo sapiens cDNA clone IMAGE:3458830 5'	601176865F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:3531953 5'	601072594F1 NIH_MGC_12 Homo sapiens cDNA clone IMAGE:3458830 5'	xz82h12.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:2871719 3'	601509696F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3911303 5'	601509696F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3911303 5'	tu18b02.x1 NCI_CGAP_Pr28 Homo sapiens cDNA clone IMAGE:2251371 3'	601302333F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3638753 5	EST177232 Jurkat T-cells VI Homo sapiens cDNA 5' end	Homo saplens chromosome 21 segment HS21,0003
Top Hit Database Source	NT	EST HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	TN	EST_HUMAN	FX	EST_HUMAN	EST_HUMAN	NT	NT	NT	TN	۲	NT	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	FN
Top Hit Acession No.	2.0E-85 AL163284.2	1760820.1	3E794306.1	3E618392.1	1.0E-85 BE618392.1	1.0E-85 BE257917.1	1.0E-85 AA778785.1	1.0E-85 AA778785.1	1.0E-85 BF311552.1	1.0E-85 BF311552.1	11417862 NT	11417862 NT	BE274217.1	7662247	7.0E-86 AA860801.1	7.0E-86 AA860801.1	11421737 NT	.38557.1	5453897 NT	11526307 NT	11417012 NT	11417012 NT	4505492 NT	BE547173.1	3E295843.1	3E547173.1	3.0E-86 AW340946.1	3E886479.1	3.0E-86 BE886479.1	N659240.1	3.0E-86 BE410354.1	2.0E-86 AA306264.1	4L163203.2
Most Similar (Top) Hit BLAST E Value	2.0E-85	2.0E-85 A	1.0E-85 Bi	1.0E-85 B	1.0E-85	1.0E-85	1.0E-85	1.0E-85	1.0E-85	1.0E-85	1.0E-85	1.0E-85	9.0E-86	7.0E-86	7.0E-86	7.0E-86	7.0E-86	7.0E-86 L3	7.0E-86	7.0E-86	7.0E-86	7.0E-86	6.0E-86	4.0E-86	4.0E-86 B	4.0E-86	3.0E-88	3.0€-88	3.0E-86	3.0E-86	3.0E-86	2.0E-86	2.0E-86
Expression Signal	1.3	1.28	244	8.09	8.09	2.06	2.56	2.56	2.28	2.28	2.7	3.43	7.66	1.3	1.33	1.33	6.91	2.91	1.83	2.35	2.15	2.15	9.33	1.48	10.18	1.9	6.02	3.31	3.31	9.01	2.02	1.33	1.67
ORF SEQ ID NO:	24489	27408		22121	22122	27704		28420	28492	28493	25283	25283		20001	20691		24865			27696	28465	28456	21027	19993	25686	19993		27963	27964	28125		20046	
SEQ ID NO:	14705	17209	12125	12224	12224	17483	18175	18175	18242	18242	19082	19082	11315	10190	10845	10845	15121	16955	17444	17475	18208	18206	11179	10175	15586	10175	15355	17718	17718	17883	19560	10231	10354
Probe SEQ ID NO:	4822	7341	2241	2344	2344	7632	8286	8236	8365	8365	9184	9448	1410	82	921	921	6164	8/0/	7593	7624	8328	8328	1272	204	2677	8567	5435	7868	7868	8734	9163	266	408

Page 299 of 413 Table 4 Single Exon Probes Expressed in Heart

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Top Hit Descriptor	yz19a08.r1 Soares_multiple_sclerosis_2NbHMSP Homo sapiens cDNA clone IMAGE:283478 5'	Human endogenous retrovirus, complete genome	Homo sapiens mRNA for KIAA1277 protein, partial cds	EST378215 MAGE resequences, MAGI Homb sapiens cDNA	Homo sapiens lysophosphatidic acid acyltransferase-delta (LPAAT-delta) mRNA, complete cds	Homo sapiens Iysophosphatitdic acid acyltransferase-delta (LPAAT-delta) mRNA, complete cds	hd87g08.x1 NCI_CGAP_GC6 Homo saplens cDNA clone IMAGE:2916542 3'	Homo sapiens cAMP-specific phosphodiesterase 8A (PDE8A) mRNA, partial cds	H.seplens mRNA encoding phospholipase c	H.saplens mRNA encoding phospholipase c	Homo sapiens butyrobetaine (gamma), 2-oxoglutarate dioxygenase (gamma-butyrobetaine hydroxylase) (BBOX), mRNA	Homo saplens butyrobetaine (gamma), 2-oxoglutarate dioxygenase (gamme-butyrobetaine hydroxylase)	(DDUCA), IIINIYA	Homo sapiens chromosome segregation 1 (Yeast nomolog) Hine (CSE LL), Illinium	Homo sapiens basic-helix-loop-helix-PAS protain (NPAS3), mRNA	Homo sapiens basic-helix-loop-helix-PAS protein (NPAS3), mRNA	Homo sapiens ribosomal protein S6 kinase, 90kD, polypeptide 5 (RPS6KA5) mRNA	Homo sapiens thyroid autoantigen 70kD (Ku antigen) (G22P1), mRNA	Homo sapiens gene for AF-6, complete cds	Homo sapiens adrencagic, beta, receptor kinase 2 (ADRBK2), mRNA	Homo sapiens NADH dehydrogenase (ubiquirione) Fe-S protein 1 (75kD) (NADH-coenzyme Q reductase) (NDUFS1) mRNA	Homo sapiens fibulin 5 (FBLN5) mRNA	Human gamma-glutamyl transpeptidase mRN/A, complete cds	Homo sapiens chromosome 21 segment HS2; C009	Homo sapiens chromosome 21 segment HS21 0009	Homo sapiens hypothetical protein (LOC51318), mRNA	Homo saplens hypothetical protein (LOC51318), mRNA	Homo saplens chromosome 21 segment HS21 C100	Homo sapiens FK506-binding protein FKB23 isoform mRNA, complete cds	Homo sapiens chromosome 21 segment HS21 C084	qb77c09.x1 Soares_fetal_heart_NbHH19W Homo saplens cDNA clone IMAGE:1706128 3' similar to SW:K1C_L_MOUSE P02535 KERATIN, TYPE I CYTOSKELETAL 10;
Top Hit Database Source	T_HUMAN			T_HUMAN		NT	EST_HUMAN	NT	IN	NT	LΝ	į	Z !			LN	LN	LN	TN	NT	L	NT	Ę	N	-N	FZ	TN	Z.	M	MT	EST_HUMAN
Top Hit Acession No.	158977.1	9635487 NT	2.0E-86 AB033103.1	2.0E-86 AW966142.1	2.0E-86 AF156776.1		2.0E-86 AW515742.1	2.0E-86 AF056480.1	216411.1	216411.1	11437135 NT	000000	1143/135 IN	11422084 NT	11545846 NT	11545846 NT	4759051 NT	11418189 NT	AB011399.1	11417883 NT	4826855 NT		20492.1	AL163209.2	AL163209.2	7706161 NT	7706181 NT	AL163300.2	AF100751.1	AL163284.2	A1150703.1
Most Similar (Top) Hit BLAST E Value	2.0E-86 N	2.0E-86	2.0E-86 /	2.0E-86	2.0E-86	2.0E-86	2.0E-86	2.0E-88	2.0E-88	2.0E-86 Z16411.1	2.0E-86	1	2.0E-88	2.0E-86	2.0E-86	2.0E-86	2.0E-86	2.0E-86	2.0E-86	2.0E-86	1.05-88	1.0E-86	1.0E-86	1.05-86	1.05-86	1.0E-86	1.0E-86	1.0E-86	1.0E-86	1.0E-86	9.0E-87
Expression Signal	2.18	2.37	1.12	1.43	2.16	2.16	2.42	2.8	1.53	1.53	2.22		2,22	1.95	2.69	2.69	1.83	2.4	1.81	1.47	3.1	2.08	2.42	1.32	1.32	11.48	11.48	6.76	1.11	2.15	1.47
ORF SEQ ID NO:	20929	21930	22011	23087				24360	25571		27057			27446	28061	28062				25139					22978				L		
Exon SEQ ID NO:	11085	12032	{	13288	١_	1		14565	15495	15495	16864	<u> </u>	_\	17241	17819	17819	ì	١.	1	19520	11483		1	1_	13179	<u> </u>	L	<u> </u>	L	L	ļ ļ
Probe SEQ ID NO:	1173	2144	2222	3369	3688	3686	3954	4679	5580	2580	6987		6987	7372	7969	7869	8276	9096	9772	9955	1570	3125	3197	3256	3258	3864	3864	4167	4832	8882	5287

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Single Exon Probes Expressed in Heart

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
6403	16264	78424	2	9.0E-87	475721 NT	LN TN	Homo saplens a disintegrin and metalloproteinase domain 22 (ADAM22), mRNA
6403	16264	26425	2	9.0E-87	4757721 NT	Ę	Homo sapiens a disintegrin and metalloproteinase domain 22 (ADAM22), mRNA
1471	10414	20233	14.17	8.0E-87	8.0E-87 X62245.1	NT	O.cuniculus mRNA for elongation factor 1 alpha
2250			2.74	7.0E-87	7.0E-87 BF063211.1	EST_HUMAN	7h86f02.x1 NCI_CGAP_Co16 Homo septens cDNA clone IMAGE:3322779 3'
2250	L		2.74	7.0E-87	7.0E-87 BF063211.1	EST_HUMAN	7h85f02.x1 NCI_CGAP_Co16 Homo saplens cDNA clone IMAGE:3322779 3'
6763	16642	26829	2.67	7.0E-87	7.0E-87 BF352776.1	EST_HUMAN	IL3-HT0619-060700-198-D10 HT0619 Homo sapiens cDNA
7794	17644	27877	3.38	7.0E-87	7.0E-87 AL043314.2	EST_HUMAN	DKFZp434N0323_r1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434N0323 5'
7794		27878	3.38	7.0E-87 AL	AL043314.2	EST_HUMAN	DKFZp434N0323_r1 434 (synonym: htes3) Homo saplens cDNA clone DKFZp434N0323 5'
8264	18144	28384	10.88	7.0E-87	7.0E-87 K03002.1	NT	Human mRNA from chromosome 15 gene with homology to MHC-HLA-SB-1 intron A
8264	18144	28385	10.88	7.0E-87	K03002.1	N	Human mRNA from chromosome 15 gene with homology to MHC-HLA-SB-1 intron A
3482		23203	0.87	6.0E-87	7657213 NT	NT	Homo sapiens hormonally upregulated neu tunior-associated kinase (HUNK), mRNA
5868	15774	25893	1.73	6.0E-87	AB029004.1	NT	Homo saplens mRNA for KIAA1081 protein, partial cds
8107	17897		3.52	6.0E-87	11432444	LΝ	Homo sepiens similar to SET translocation (myeloid leukemia-associated) (H. sapiens) (LOC63102), mRNA
1142	11056	L	1.69	5.0E-87	5.0E-87 AA382811.1	EST_HUMAN	EST86094 Testis I Homo saplens cDNA 5' end
9450	11056	20898	1.53	5.0E-87	5.0E-87 AA382811.1	EST_HUMAN	EST96094 Testis I Homo sapiens cDNA 5' end
950	10874	20721	1.33	4.0E-87		N	Homo sapiens chromosome 21 segment HS21C010
1165	11068	20912	10.54	4.0E-87	4.0E-87 AB037835.1	NT	Homo sapiens mRNA for KIAA1414 protein, pertial cds
1411	11316	21179	0.86	4.0E-87	4.0E-87 R78133.1	EST HUMAN	yi80f10.r1 Soares placenta Nb2HP Homo sapiens cDNA clone IMAGE:145579 5' similar to contains Alurepetitive element.
1985	11878	21771	0.92	4.0E-87	7.	LN	Homo sapiens mRNA for KIAA0456 protein, partial cds
2372	12252	22142	1.07	4.0E-87	7706299 NT	NT	Homo sapiens CGI-60 protein (LOC51626), mRNA
2372		22143	1.07	4.0E-87	TV06299 NT	FN	Homo saplens CGI-60 protein (LOC51626), mRNA
3419	13338	23140	2.19	4.0E-87	5174574 NT	LN	Homo sapiens mysloid/lymphoid or mixed-lineage leukemia (trithorax (Drosophila) homolog); translocated to, 4
6343	15264	26090	6.47	4.0E-87 000321	000321	SWISSPROT	ETS-RELATED PROTEIN 71 (ETS TRANSLOCATION VARIANT 2)
5686	15595	25696	4.36	4.0E-87 BE	BE247284.1	EST_HUMAN	TCBAP1E4051 Pediatric pre-B cell acute lymphoblastic leukemia Baylor-HGSC project=TCBA Homo sapiens, cDNA clone TCBAP4051
8505	18378	28644	4.35	4.0E-87	M60676.1	N	Human von Willebrand factor pseudogene corresponding to exons 23 through 34
8994	18798		2.13	4.0E-87	11417339 NT	TN	Homo sapiens similar to heat shock 70kD protein 9B (mortalin-2) (H. sapiens) (LOC63184), mRNA
9692	1		14.77	4.0E-87	11417812 NT	TN	Homo sapiens purinergic receptor P2X-like 1, orphan receptor (P2RXL1), mRNA
2749		22502	2.99	2.0E-87	4885420	NT	Homo sapiens high-mobility group (nonhistone chromosomal) protein 4 (HMG4) mRNA
3717	$_{\perp}$	23414	0.89	2.0E-87 AL	J116935.1	EST HUMAN	AU116935 HEMBA1 Homo sepiens cDNA clone HEMBA1000307 5
4826	14708	24492	1.17	2.0E-87 BF	BF376311.1	EST_HUMAN	CM0-TN0038-150900-552-h08 TN0038 Homo sapiens cDNA

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Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
4880	14760	24537	0.8	2.0E-87	BE175478.1	EST_HUMAN	RC5-HT0580-200300-031-G04 HT0580 Homo saplens cDNA
5473	15393	25457	78.7	2.0E-87	BE734190.1	EST_HUMAN	601569041F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3843730 5'
5473	15393	25458	7.87	2.0E-87	BE734190.1	EST_HUMAN	601569041F1 NIH_MGC_21 Homo saplens cDNA clone IMAGE:3843730 5'
5819	16726		6.27	2.0E-87	BE567193.1	EST_HUMAN	601341383F1 NIH_MGC_53 Homo sapiens cDNA clone IMAGE:3683348 5'
9529	16122	28275	1.51	2.0E-87	BE294432.1	EST_HUMAN	601176032F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:3531511 5'
6407	16268	26430	32.21	2.0E-87		EST_HUMAN	yv21e07.r1 Soares fetal liver spleen 1NFLS Homo saplens cDNA clone IMAGE:243396 5'
6526	16385	26564	28.31	2.0E-87	N48128.1	EST_HUMAN	yv21e07.r1 Soares fetal liver spleen 1NFLS Homo saplens cDNA clone IMAGE:243396 5
6892	16771	26966	3.81	2.0E-87	X52851.1	NT	Human cyclophilin gene for cyclophilin (EC 5.2 1.8)
7634	17485		5.11	2.0E-87	BE531136.1	EST_HUMAN	601278315F1 NIH_MGC_39 Homo sapiens cDNA clone IMAGE:3610539 5'
1165	12845		2.33	1.0E-87	7705683 NT	NT	Homo sapiens putative glycolipid transfer protein (LOC51054), mRNA
1413	11318	21181	0.94	1.0E-87	AW361977.1	EST_HUMAN	PM2-CT0265-141099-001-904 CT0265 Homo sapiens cDNA
1413	11318	21182	0.94	1.0E-87	AW361977.1	EST_HUMAN	PM2-CT0265-141099-001-g04 CT0265 Homo sapiens cDNA
3849	13563	23349	3.18	1.0E-87	Y00052.1	NT	Human mRNA for T-cell cyclophilin
3873	13587	23374	2.47	1.0E-87	4768827 NT	TN	Homo sapiens neurexin III (NRXN3) mRNA
2021	14927	24699	1.04	1.0E-87	U50949.1	NT	Rattus norvegicus taste bud receptor protein TB 641 (TB 641) gene, complete cds
5774	15681	25788	3.39	1.0E-87		NT	Homo sapiens growth factor receptor-bound protein 10 (GRB10) gene, exon 8
5774	15681	26789	3.39	1.0E-87	AF073371.1	NT	Homo saplens growth factor receptor-bound protein 10 (GRB10) gene, exon 8
6378	16238	26398	1.6	1.0E-87	11431590 NT	NT	Homo saplens protein kinase C, beta 1 (PRKCB1), mRNA
6724	16604	26793	13.13	1.0E-87	AF214562.1	NT	Homo sapiens tracheal epithellum enriched protein (PLUNC) gene, complete cds
7165	17042	27233	1.19	1.0E-87,	AB022918.1	NT	Homo sapiens mRNA for alpha2,3-sialyltransferase ST3Gal VI, complete cds
7165	17042		1.19	1.0E-87		NT	Homo sapiens mRNA for alpha2,3-slalyltransferase ST3Gal VI, complete cds
7554	17405	27820	2.77	1.0E-87		EST_HUMAN	RC6-BN0276-050700-012-E02 BN0276 Homo saplens cDNA
7554	17405	27621	2.77	1.0E-87	BE818183.1	EST_HUMAN	RC8-BN0276-050700-012-E02 BN0276 Homo sapiens cDNA
8114	18003	28249	2.08	1.0E-87	5729867 NT	NT TA	Homo sapiens hect domain and RLD 2 (HERC2), mRNA
8367	18244		1.78	1.0E-87	D10083.1	NT	Homo sapiens RGH1 gene, retrovirus-like element
6236	19751		2.02	1.0E-87	T657632 NT	NT	Homo sapiens sulfotransferase-related protein (SULTX3), mRNA
1090	11008	20847	7.39	9.0E-88	AF167465.1	NT	Homo sapiens double strended RNA activated protein kinase (PKR) gene, exon 12
1327	11234	21090	2	9.0E-88	AB037820.1	NT	Homo sapiens mRNA for KIAA1399 protein, pertial cds
1327	11234	11091	2	9.0E-88		NT	Homo sapiens mRNA for KIAA1399 protein, partial cds
3574	13488	23279	1.13	9.0E-88	AL163209.2	NT	Homo sapiens chromosome 21 segment HS210009
4172	14072		2.64	9.0E-88	X91929.1	NT	H.sapiens ECE-1 gene (exon 9)
4172	14072	23848	2.64	9.0E-88	X91929.1	NT	H.sapiens ECE-1 gene (exon 9)
4834	14812	24580	1.23	9.0E-88	AB026898.1	TN	Homo saplens DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes, complete cds)
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Top Hit Descriptor	Homo sepiens X-linked anhidroitic ectodermei dysplasia protein gene (EDA), exon 2 and flanking repeat regions	Homo sapiens KIAA0063 gane product (KIAA0063), mRNA	K9719F Human fetal heart, Lambda ZAP Express Homo sapiens cDNA clone K9719 6' similar to ZINC FINGER PROTEIN HZF1	Homo sapiens intersectin short isoform (ITSN) mRNA, complete cds	Homo saplens intersectin short isoform (ITSN) mRNA, complete cds	Homo sapiens intersectin short isoform (ITSN) mRNA, complete cds	wd68h08.x1 NCI_CGAP_Lu24 Homo sapienis cDNA clone IMAGE:2336799 3' similar to contains Alu repetitive element,contains element MER22 MER22 repetitive element;	Homo sapiens intersectin short isoform (ITSN) mRNA, complete cds	Homo sapiens intersectin short isoform (ITSN) mRNA, complete cds	ym06b10.r1 Soares Infant brain 1NIB Homo sapiens cDNA clone iMAGE:47129 5	Homo sapiens chromosome 21 segment HS210084	PM1-TN0028-050900-004-f10 TN0028 Homp sapiens cDNA	PM1-TN0028-050900-004-f10 TN0028 Hamp sapiens cDNA	Homo sapiens transforming growth factor, beta-induced, 68kD (TGFBI), mRNA	Homo sapiens cell division cycle 10 (homologicus to CDC10 of S. cerevisiae) (CDC10) mRNA	Homo saplens KIAA0152 gene product (KIAA0152), mRNA	Homo sapiens KIAA0152 gene product (KIAA0152), mRNA	Homo sapiens hypothetical protein FLJ21634 (FLJ21634), mRNA	Homo saplens zinc finger protein 259 (ZNF259) mRNA	za48f12.s1 Soares fetal liver spleen 1NFLS Homo capiens cDNA clone IMAGE:295823 3'	Homo saplens a disintegrin and metalloproteir ase domain 23 (ADAM23) mRNA	Homo saplens a disintegrin and metalloprotetriase domain 23 (ADAM23) mRNA	Homo sapiens hypothetical protein FLJ20220 (FLJ20220), mRNA	Homo saplens valosin-containing protein (VCP), mRNA	Homo sapiens polycythemia rubra vera 1; cell surface receptor (PRV1), mRNA	Homo sapiens v-ral simian leukemia viral onccigene homolog A (ras related) (RALA), mRNA	Homo saplens putative anion transporter 1 mRNA, complete cds	Homo sapiens retinoblastoma-binding protein 2 (RBBP2), mRNA	Homo sapiens growth differentiation factor 5 (cartilage-derived morphogenetic protein-1) (GDF5), mRNA
Top Hit Database Source	Ę	N	EST HUMAN	Į.	F	۲	EST HUMAN	F	F	EST HUMAN	Ę	EST HUMAN	EST HUMAN	F	Ę	NT.	IN.	F	F	EST_HUMAN	NT	NT	NT	TN	TN	IN	Ę	NT	LN.
Top Hit Acession No.	AF003528.1	7661887 NT	N89399.1	AF114488.1	AF114488.1	AF114488.1	AI693217.1	AF114488.1	AF114488.1	H10932.1	AL163284.2	BF091229.1	BF091229.1	11416585 NT	4502694 NT	7661947 NT	7661947 NT	11545800 NT	4508020 NT	N66951.1	4501912 NT	4501912 NT	11429300 NT	11429567 NT	TN 8888888	11420697 NT	AF279265.1	11436400 NT	11421728 NT
Most Similar (Top) Hit BLAST E Value	6.0E-88	-	5.0E-88	5.0E-88	5.0E-88	5.0E-88	5.0E-88	5.0E-88			5.0E-88	4.0E-88	4.0E-88	4.0E-88	4.0E-88	4.0E-88	4.0E-88	3.0E-88	3.0E-88	3.0E-88	3.0E-88	3.0E-88	3.0E-88	3.0E-88	3.0E-88	3.0E-88	3.0E-88	3.0E-88	3.0E-88
Expression Signal	3.69	1.19	8	0.92	0.94	0.94	2.31	0.91	0.87	264	1.84	1.64	1.64	1.93	284	2.1	2.1	0.83	4.77	4.31	1.21	1.21	3.17	4.09	3.84	3.39	12.03	6.66	8.62
ORF SEQ ID NO:	27284		22368		22704			23212	Ĺ	26185	26694	21070	21071	26316	28402		28884	L			23819	23820		24937	25403	25480	26098	28479	26688
Exan SEQ ID NO:	17093	11685	12470	12897	12905	12905	13263	13407	14519	16042	16507	11214	11214	16159	18160	18595	18595	10647	11669	12845	14047	14047	14276	15166	15349	15417	15984	18313	16500
Probe SEQ ID NO:	7216	1787	2602	2970	2978	2978	3343	3491	4631	6909	6627	1307	1307	6295	8281	8780	8780	715	1770	2918	4147	1414	4380	5242	5429	5498	6204	6452	6620

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Single Exon Probes Expressed in Heart

Homo saplens molybdenum cofactor blosynthesis protein A and molybdenum cofactor biosynthesis protein C aa54a11.s1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:824732 3' similar to WP:B0272.2 os91g03.s1 NCI_CGAP_GC3 Homo septens oDNA clone IMAGE:1612766 3' similar to gb:M16342 HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEINS C1/C2 (HUMAN); (CSEN) mRNA UI-H-BI1-asa-d-04-0-UI.s1 NCI_CGAP_Sub3 Homo sapiens cDNA clone IMAGE:2718750 3'
UI-H-BI1-asa-d-04-0-UI.s1 NCI_CGAP_Sub3 Homo sapiens cDNA clone IMAGE:2718750 3'
Homo sapiens KIAA0417 mRNA, complete cds
Homo sapiens KIAA0417 mRNA, complete cds DKFZp434N0323_r1 434 (synonym: htes3) Homo sepiens cDNA clane DKFZp434N0323 5 DKFZp434E246_r1 434 (synonym: hies3) Homo saplens cDNA clone DKFZp434E246 5 Homo saplens actin related protein 2/3 complex, subunit 1A (41 kD) (ARPC1A), mRNA Homo sapiens v-ets avian erythroblastosis virus E26 oncogene related (ERG), mRNA Homo sapiens hormonally upregulated neu tumor-associated kinase (HUNK), mRNA Homo sapiens hormonally upregulated neu turnor-associated kinase (HUNK), mRNA Homo sapiens Calsenilin, presenllin-binding protein, EF hand transcription factor (Homo saplens Inner membrane protein, mitochondrial (mitofilin) (IMMT), mRNA Homo sapiens serine/threonine-protein kinase PRP4 homolog (PRP4) mRNA 601142409F1 NIH_MGC_14 Homo sapiens cDNA clone IMAGE:3506186 5 Homo sapiens complement component 8, beta polypeptide (C8B) mRNA Homo sapiens SNARE protein kinase SNAK inRNA, complete cds Homo sapiens dyneln, axonemal, light polypeptide 4 (DNAL4), mRNA Homo sapiens transcobalamin II; macrocytic enemia (TCN2), mRNA Homo sapiens SNARE protein kinase SNAK inRNA, complete cds Top Hit Descriptor Homo saplens mRNA for KIAA0823 protein, partial cds Homo saplens mRNA for KIAA0823 protein, partial cds Homo sapiens chromosome 21 segment HS21C046 Human aconitate hydratase (ACO2) gene, exon 2 Homo saptens transgelin 2 (TAGLN2), mR H.sapiens CLN3 gene, complete CDS H.sapiens CLN3 gene, complete CDS H.sapiens Wee1 hu gene H.sapiens Wee1 hu gene mRNA, complete cds CE00851; **EST HUMAN** HUMAN EST_HUMAN EST_HUMAN **EST HUMAN** EST_HUMAN EST_HUMAN Top Hit Database Source EST 눋 눋 눋 눋 11420754 NT z 눋 뉟 4506124 NT F 눋 7857213 NT 눋 z 'n 11417974 NT 5803114 7657213 7305198 11421238 5031668 4557390 **Top Hit Acession** 2.0E-88 7305 2.0E-88 AF246219.1 2.0E-88 AF246219.1 2.0E-88 503 1.0E-88 AW139565.1 .0E-88 AW 139565.1 AB020630.1 1.0E-88 AA488981.1 BE311557.1 AL043314.2 1.0E-88 AA991479.1 7.0E-89 AL045748.1 3.0E-88 AF034374.1 AB020630. .0E-88 AB007877. ġ 7.0E-89 X62048.1 X99832.1 X99832.1 7.0E-89 U87927.1 7.0E-89 7.0E-89 .0E-88 8.0E-89 7.0E-89 7.0E-89 1.0E-88 1.0E-88 7.0E-89 7.0E-89 3.0E-88 3.0E-88 7.0E-89 7.0E-89 6.0E-89 9.0E-89 dost Similar (Top) Hit BLAST E Value 4.12 5.14 1.35 1.35 1.35 2.94 1.12 1.42 4.19 1.93 4.98 4.98 17.59 4.06 2.95 2.27 1.35 2.12 Expression Signal 28098 28108 20768 21954 26836 21486 24007 25597 25598 20194 24457 25077 26450 28097 28107 26626 20780 26033 26282 20195 24511 21372 28135 28447 ORF SEQ ÖΝΘ 17856 17862 10924 15518 15518 16128 17891 18198 14870 15254 17856 12053 16440 10938 11512 11617 14225 15909 15909 10371 14728 16289 19117 10371 SEQ ID 15254 ÿ 8008 8012 8012 8742 4785 5334 8008 2188 1716 4328 5604 5604 6004 6004 6263 7524 6428 1006 6269 4847 9920 Probe SEQ ID 1020 9502 8321 426 426 5334 7427 9286 ÿ

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Single Exon Probes Expressed in Heart

r CBAP2E0383 Pediatric pre-B cell acute lymohoblastic leukemia Baylor-HGSC project=TCBA Homo sapiens CBAP2E0383 Pediatric pre-B cell acute lymphoblastic leukemia Baylor-HGSC project=TCBA Homo sapiens Homo saplens solute carrier family 24 (sodium/potassium/calcium exchanger), member 2 (SLC24A2), mRNA qg98c08.x1 Soares_NFL_T_GBC_S1 Homo capiens cDNA clone IMAGE:1843022 3' similar to gb:J04131 GAMMA-GLUTAMYLTRANSPEPTIDASE_1 PRECURSOR (HUMAN);contains Alu repetitive element; Homo sapiens cell adhesion molecule with homology to L1CAM (close homologue of L1) (CHL1), mRNA Human MAGE-7 anigen (MAGE7) pseudogerie, complete cds Homo sapiens integrin, alpha 3 (antigen CD49C, alpha 3 subunit of VLA-3 receptor) (ITGA3), mRNA Homo sapiens topoisomerase-related function protein (TRF4-2) mRNA, partial cds Homo sapiens ubiquitin-conjugating enzyme E2L 3 (UBE2L3) mRNA Homo sapiens HSPC159 protein (HSPC159); mRNA Homo sapiens ubiquitin-conjugating enzyme E2L 3 (UBE2L3) mRNA **Top Hit Descriptor** EST388290 MAGE resequences, MAGN Homo sapiens cDNA cDNA clone TCBAP0383 QV3-NT0022-080600-219-g03 NT0022 Homo sapiens cDNA H.sapiens HCK gene for tyrosine kinase (PTK), exons 10-11 H.saplens HCK gane for tyrosine kinase (PTK), exons 10-11 partial cds AV705749 ADB Homo sapiens cDNA clone ADBBGA01 5' Homo sapiens mRNA for KIAA0406 protein, partial cds Homo sapiens mRNA for KIAA0406 proteln, partial cds Homo sapiens mRNA for KIAA1342 protein, partial cds Human N-ethylmaleimide-sensitive factor mRNA, Homo sapiens PXR2b protein (PXR2b), mRNA Homo sapiens PXR2b protein (PXR2b), mRNA Homo sapiens PXR2b protein (PXR2b), mRNA Homo sapiens PXR2b protein (PXR2b), mRNA Homo sapiens gene for LECT2, complete cds Human GT24 (GT24) mRNA, partial cds Homo septens GGT gene, exon 5 cDNA clone TCBAP0383 EST_HUMAN EST_HUMAN EST_HUMAN HUMAN EST_HUMAN EST_HUMAN Top Hit Database Source EST 토토토토토 11434411 NT 11433673 NT 눋 11428801 NT 눋 눋 7706670 NT 뉟 4507788 7661817 7706670 4507788 **Fop Hit Acession** 4.0E-89 BE762749.1 3.0E-89 AW976181.1 2.0E-89 AF089897.1 2.0E-89 X58742.1 2.0E-89 X58742.1 AB007866.2 AB007866.2 5.0E-89 BE244323.1 5.0E-89 BE244323.1 AB037763.1 BE762749.1 AV705749.1 2.0E-89 AI222095.1 2.0E-89 AJ007378.1 2.0E-89 AB007546. ģ U03985.1 **U81004.1** 2.0E-89 U10692.1 2.0E-89 2.0E-89 t 6.0E-89/ 3.0E-89 / 2.0E-89 2.0E-89 2.0E-89 2.0E-89 6.0E-89 6.0E-89 6.0E-89 2.0E-89 (Top) Hit BLAST E Aost Similar Value 6.18 1.92 1.46 1.14 0.91 0.91 1.45 6.18 1.6 4.93 3.73 2.83 4.87 2.24 1.97 0.84 3.5 2.77 2.77 <u>1</u>9 Expression Signal 23741 24076 25123 26699 28953 29055 22563 25215 20175 20175 20178 23740 25526 28557 28851 22154 22155 20176 ORF SEQ ID NO: 24646 24647 23201 24212 24211 18666 10348 10348 13963 16510 18568 10348 10348 12780 13963 14292 15288 18762 12263 12263 16336 12775 15455 16379 13396 14430 14430 19241 SEQ ID 14881 14881 2383 402 402 519 2852 4053 4061 4061 5368 6538 6520 888 8854 8955 Probe SEQ ID 2847 9688 6630 3480 4537 5007 5007 4537 421 121

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Single Exon Probes Expressed in Heart

			-	,	_		_	_	_	_	_	_			<u>~~</u>	~***		-	_==	~~	-		-	~	-	<u> </u>		7 /4-	
Top Hit Descriptor	hr81d09.x1 NCI_CGAP_KId11 Homo saplens cDNA clone IMAGE:3134897 3' similar to TR:054778 054778 SOLUTE CARRIER FAMILY 22 -LIKE 2 PROTEIN;	hr81d09.x1 NCI_CGAP_KId11 Homo sapiens dONA clone IMAGE:3134897 3' similar to TR:054778 054778 SCLUTE CARRIER FAMILY 22 -LIKE 2 PROTEIN ;	Homo sapiens chromosome 21 segment HS21C046	Hamo sapiens chromosome 21 segment HS21C046	Homo sapiens chromosome 21 segment HS21C046	Homo sapiens chromosome 21 segment HS21C046	7e36f08.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3284583 3'	7e36f08.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3284583 3'	Homo sapiens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively spliced	ai63d08.s1 Soares_testis_NHT Homo sapiens cDNA clone 1375503 3'	601655837R1 NIH_MGC_66 Homo sapiens cDNA clone IMAGE:3855824 3'	601655837R1 NIH_MGC_66 Homo saplens cONA clone IMAGE:3855824 3'	y86e04.s1 Soares fetal liver spleen 1NFLS Homo saplens cDNA clone IMAGE:212190 3' similar to SP:C1TC_HUMAN P11586 C-1-TETRAHYDROFOLATE SYNTHASE, CYTOPLASMIC;	y86e04.s1 Soares fetal liver spleen 1NFLS Homo septens cDNA clone IMAGE:212190 3' similar to SP:C1TC HUMAN P11886 C-1-TETRAHYDROFOLATE SYNTHASE, CYTOPLASMIC.	H.sapiens ECE-1 gene (exon 6)	H.sapiens ECE-1 gene (exon 6)	Homo sapiens hypothetical protein FLJ10388 (FLJ10388), mRNA	Homo saplens hypothetical protein FLJ10388 (FLJ10388), mRNA	Homo sapiens HsGCN1 mRNA, partial cds	Homo sepiens HsGCN1 mRNA, partial cds	Homo saplens inositol 1,4,5-triphosphate receptor, type 3 (ITPR3) mRNA	Homo sapiens inositol 1,4,5-triphosphate receptor, type 3 (ITPR3) mRNA	Homo sapiens TOL6 gene, exon 1-10b	Human gamma-aminobutyric acid transaminase mRNA, partial cds	Homo sapiens intersectin long isoform (ITSN) mRNA, complete cds	Homo saplens pregnancy-zone protein (PZP) inRNA	z/82g10.s1 Soares_fetal_liver_spleen_1NFLS1S1 Homo sapiens cDNA clone IMAGE:461442 3'	zj82g10.s1 Soares_fetal_liver_spleen_1NFLS_S1 Homo saplens cDNA clone IMAGE:461442 3'	DKFZp762P1616_r1 762 (synonym: hmel2) Hömo sapiens cDNA clone DKFZp762P1616 5'
Top Hit Database Source	EST_HUMAN	EST_HUMAN	NT	LN	NT	ΙN	EST_HUMAN	EST_HUMAN	LN	EST_HUMAN	EST HUMAN	EST_HUMAN	EST HUMAN	EST HUMAN	NT	LN	Z	TN	₽N	NT TA	L	Z	NT	LΝ	NT	NT	EST_HUMAN	EST_HUMAN	EST HUMAN
Top Hit Acession No.	BF196052.1	BF198052.1	AL163246.2	AL163246.2	AL163246.2	AL163246.2	BE670561.1	BE670561.1	AF223391.1	AA782977.1	BE962525.2	BE962525.2	H68849.1	H68849.1	X91926.1	X91926.1	TN 8922398	8922398 NT	U777700.1	U77700.1	4504794 NT	4504794 NT	AB035344.1	U80226.1	AF114487.1	4506354 NT	AA705222.1	AA705222.1	AL135549.1
Most Similar (Top) Hit BLAST E Value	1.0E-89	1.0E-89	9.0E-90	9.0E-80	8.0E-90	8.0E-90	8.0E-90	8.0E-90	7.0E-90	7.0E-90	7.0E-90	_	7.05-90			6.0E-90	6.0E-90	6.0E-90	6.0E-90	6.0E-90	6.0E-90	6.0E-90	5.0E-90	5.0E-90	5.0E-80	5.0E-90	5.0E-90	5.0E-80	6.0E-90
Expression Signal	6.12	6.12	1.16	1.16	1.62	2.93	5.28	5.28	2.65	1.91	1.88	1.88	1.98	1.98	1.14	1.14	7.33	7.33	3.64	3.54	3.26	3.26	10.5	1.55	2.19	3.08	1.07	1.07	0.98
ORF SEQ ID NO:	28960	28961	26859	26860	20808	20806	21072	21073			27260		27927			22750	L	23810	25646	25647		L			22273		24186	24187	24251
Exan SEQ ID NO:	18672	18672	16668	16668	10965	10965	12690	12690	10748	16785	L_	_	17683	17683	12957	12957	14034	14034	15564	15554		ļ	10123	11087	12382	14334	14400	14400	14463
Probe SEQ ID NO:	0988	988	62.83	6789	1047	1048	1308	1308	818	2089	7183	7183	7833	7833	3029	3028	4134	4134	6841	5641	6846	6846	149	1175	2508	4440	4507	4607	4571

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								sterase 3 (H. sapiens) (LOC63214),	sterase 3 (H. saplens) (LOC63214). ype 8A, member 2 (ATP8A2), mRNA	isterase 3 (H. sapiens) (LOC63214). ype 8A, member 2 (ATP8A2), mRNA	isterase 3 (H. sapiens) (LOC63214), ype 8A, member 2 (ATP8A2), mRNA GE:2128761 3'	sterase 3 (H. sapiens) (LOC63214), ype 8A, member 2 (ATP8A2), mRNA GE:2128761 3'	sterase 3 (H. sapiens) (LOC63214), ype 8A, member 2 (ATP8A2), mRNA GE:2128761 3'	sterase 3 (H. sapiens) (LOC63214), ype 8A, member 2 (ATP8A2), mRNA GE:2128761 3'	sterase 3 (H. sapiens) (LOC63214), ype 8A, member 2 (ATP8A2), mRNA GE:2128761 3' A	sterase 3 (H. sapiens) (LOC63214), ype 8A, member 2 (ATP8A2), mRNA GE:2128761 3' A	sterase 3 (H. sapiens) (LOC63214), ype 8A, member 2 (ATP8A2), mRNA 3E:2128761 3' A	sterase 3 (H. sapiens) (LOC63214), ype 8A, member 2 (ATP8A2), mRNA 3E:2128761 3' A	sterase 3 (H. sapiens) (LOC63214), ype 8A, member 2 (ATP8A2), mRNA SE:2128761 3' A	sterase 3 (H. sapiens) (LOC63214), ype 8A, member 2 (ATP8A2), mRNA 3E:2128761 3' A A 38147 5'	sterase 3 (H. sapiens) (LOC63214), ype 8A, member 2 (ATP8A2), mRNA 3E:2128761 3' A A 17 (HMG17), mRNA	sterase 3 (H. sapiens) (LOC63214), ype 8A, member 2 (ATP8A2), mRNA GE:2128761 3' A A 17 (HMG17), mRNA 17 (HMG17), mRNA	sterase 3 (H. sapiens) (LOC63214). ype 8A, member 2 (ATP8A2), mRNA GE:2128761 3' A A 17 (HMG17), mRNA 17 (HMG17), mRNA ins cDNA clone IMAGE:1713410 3' ins cDNA clone Clone Clone Clone Clone Clone Clone Clone Clone Clone Clone Clone Clone Clone Clone Clone	sterase 3 (H. sapiens) (LOC63214), ype 8A, member 2 (ATP8A2), mRNA 3E:2128761 3' A 17 (HMG17), mRNA 17 (HMG17), mRNA ins cDNA clone IMAGE:1713410 3' IR3.;	sterase 3 (H. sapiens) (LOC63214). ype 8A, member 2 (ATP8A2), mRNA 3E:2128761 3' A 17 (HMG17), mRNA 17 (HMG17), mRNA 17 (HMG17), mRNA 183 5' Ins cDNA clone IMAGE:1713410 3' R3 .:	sterase 3 (H. sapiens) (LOC63214), ype 8A, member 2 (ATP8A2), mRNA 3E:2128761 3' A A 17 (HMG17), mRNA 17 (HMG17), mRNA ns cDNA clone IMAGE:1713410 3' R3.: R3.:	sterase 3 (H. sapiens) (LOC63214), ype 8A, member 2 (ATP8A2), mRNA 3E:2128761 3' A A 17 (HMG17), mRNA 17 (HMG17), mRNA ins cDNA clone iMAGE:1713410 3' R3.: R3.: SA) (H. sapiens) (LOC83484), mRNA	sterase 3 (H. sapiens) (LOC63214), ype 8A, member 2 (ATP8A2), mRNA 3E:2128761 3' A A 17 (HMG17), mRNA 17 (HMG17), mRNA Ins cDNA clone IMAGE:1713410 3' R3.: R3.: SA) (H. sapiens) (LOC63484), mRNA SA) (H. sapiens) (LOC63484), mRNA
. Top Hit Descriptor			nartial cds	vartial cds		RNA	RNA FLJ13222), mRNA	Homo sapiens adenylate cyclase 9 (ADCY9) mRNA Homo sapiens hypothetical protein FLJ13222 (FLJ13222), mRNA Homo sapiens similar to ectonucleotide pyrophosphatase/phosphodiestarase 3 (H. sapiens) (LOC63214), mRNA	Homo saplens adenylate cyclase 9 (ADCY9) mRNA Homo saplens hypothetical protein FLJ13222 (FLJ13222), mRNA Homo saplens similar to ectanucleotide pyropriosphatasalphosphodiesterase 3 (H. saplens) (LOG3214), mRNA Homo saplens ATPase, aminophospholipid transporter-like, Class I, type 8A, member 2 (ATP8A2), mRNA	RNA FLJ13222), mRNA osphatase/phosphodiestarase 3 (H. sapi nsporter-like, Class I, type 8A, member /	Homo sepiens adenylate cyclase 9 (ADCY9) mRNA Homo sepiens hypothetical protein FLJ13222 (FLJ13222), mRNA Homo sepiens similar to ectonucleotide pyrophiosphatase/phosphodiesterase 3 (H. sapimRNA Homo sepiens ATPase, aminophospholipid transporter-like, Class I, type 8A, member 2 Homo sepiens gene for AF-6, complete cds ar78h05.x1 Barstead aorta HPLRB6 Homo sapiens cDNA clone IMAGE:2128761 3'	RNA FLJ13222), mRNA osphatase/phosphodiesterase 3 (H. sapi nsporter-like, Class I, type 8A, member 2 irens cDNA clone IMAGE:2128761 31 A	RNA FLJ13222), mRNA osphatase/phosphodiesterase 3 (H. sapi nsporter-like, Class I, type 8A, member 2 iens cDNA clone IMAGE:2128761 3' A A	RNA FLJ13222), mRNA osphatase/phosphodiesterase 3 (H. sapl nsporter-like, Class I, type 8A, member 5 iens cDNA clone IMAGE:2128761 3' A A A A A A A A A A A A A A A A A A A	RNA FLJ13222), mRNA osphatase/phosphodiesterase 3 (H. sapl nsporter-like, Class I, type 8A, member 2 iens cDNA clone IMAGE:2128761 3' A A A nunit 1 (MYPT1), mRNA osine kinase, exon 16	RNA FLJ13222), mRNA osphatase/phosphodiesterase 3 (H. sapi nsporter-like, Class I, type 8A, member 2 iens cDNA clone IMAGE:2128761 3' A A A A A A A A A A A A A A A A A A A	RNA FL/13222), mRNA osphatase/phosphodiesterase 3 (H. sapi nsporter-like, Class I, type 8A, member 2 iens cDNA clone IMAGE:2128761 3' A A unit 1 (MYPT1), mRNA osine kinase, exon 16 sin, complete cds artial cds	RNA FL/13222), mRNA osphatase/phosphodiesterase 3 (H. sapl rsporter-like, Class I, type 8A, member 2 iens cDNA clone IMAGE:2128761 3' A A A A A A A A A A A A A A A A A A A	RNA FL/13222), mRNA osphatase/phosphodiesterase 3 (H. sapl rsporter-like, Class I, type 8A, member 2 iens cDNA clone IMAGE:2128761 3' A A A A A A A A A A A A A A A A A A A	RNA FL/13222), mRNA osphatase/phosphodiesterase 3 (H. sapi reporter-like, Class I, type 8A, member 2 iens cDNA clone IMAGE:2128761 3' A A unit 1 (MYPT1), mRNA osine kinase, exon 16 sin, complete cds artial cds Ogene, exon 8 Ona clone IMAGE:3689147 5' ONA clone IMAGE:3453834 5'	Homo sapiens adenylate cyclase 9 (ADCY9) mRNA Homo sapiens similar to ectonucleotide pyrophosphatase/phosphodiesterase 3 (H. sapiens mRNA Homo sapiens similar to ectonucleotide pyrophosphatase/phosphodiesterase 3 (H. sapiens mRNA Homo sapiens ATPase, aminophospholipid 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	H.sapiens mRNA encoding phospholipase c	H.sapiens mRNA encoding phospholipase c	Homo saplens anglopoletin 4 (ANG4) mRNA, partial cds	Homo sapiens angiopoletin 4 (ANG4) mRNA, partial cds	Homo sapiens adenylate cyclase 9 (ADCY9) mRNA	Homo sapiens hypothetical protein FLJ13222 (FLJ13222), mRNA	sociation avgoration	boudaité association	ninophospholipid trans	mRNA Homo sapiens ATPase, aminophospholipid transi Homo sapiens gene for AF-6, complete cds	ninophospholipid transi-6, complete cds HPLRB6 Homo sapie	mRNA Homo sapiens ATPase, aminophospholipid transl Homo sapiens Gene for AF-6, complete cds ar78h05.x1 Barstead aorta HPLRB6 Homo sapier Homo sapiens chromosome 21 unknown mRNA	MRNA Homo sapiens animar to eccuracione propries Homo sapiens ATPase, aminophospholipid trans Homo sapiens gene for AF-6, complete cds ar78h05.x1 Barstead aorta HPLRB6 Homo sapien Homo sapiens chromosome 21 unknown mRNA Homo sapiens chromosome 21 unknown mRNA	ninophospholipid trensi -6, complete cds HPLRB6 Homo sapiel e 21 unknown mRNA e 21 unknown mRNA sphatase, target subur	ninophospholipid trensi -6, complete cds HPLRB6 Homo sapiel e 21 unknown mRNA e 21 unknown mRNA sphatase, target subur silscoldin receptor tyros	ninophospholipid transi -6, complete cds HPLRB6 Homo sapirer e 21 unknown mRNA e 21 unknown mRNA sphatase, target subur iliscoldin receptor tyros	MRNA Homo sapiens ATPase, aminophospholipid transporter- Homo sapiens ATPase, aminophospholipid transporter- Homo sapiens gene for AF-6, complete cds ar78h05.x1 Barstead aorta HPLRB6 Homo sapiens cDN Homo sapiens chromosome 21 unknown mRNA Homo sapiens chromosome 21 unknown mRNA Homo sapiens myosin phosphatase, target subunit 1 (M H. sapiens gene encoding discoldin receptor tyrosine kin Homo sapiens DNA for amyloid precursor protein, comp	ninophospholipid trensi -6, complete cds -6, complete cds HPLRB6 Homo sapirer e 21 unknown mRNA e 21 unknown mRNA sphatase, target subur sphata	ninophospholipid trensi -6, complete cds -HPLRB6 Homo sapier e 21 unknown mRNA e 21 unknown mRNA sphatase, target subur silscoldin receptor tyros lydid precursor protein, part rding enzyme (NEC2) (ninophospholipid trensinophosp	ninophospholipid trensinophosp	ninophospholipid trensinophosp	inophospholipid trensing complete cds 4. complete cds HPLRB6 Homo sapier e 21 unknown mRNA e 21 unknown mRNA sphatase, target subun ilscoldin receptor tyros yold precursor protein (IAA1244 protein, part fing enzyme (NECZ) g 99 Homo sapiens cDN 10 Homo sapiens cDN y group (nonhistonejet y group (nonhiston	MRNA Homo sapiens Sinnia to ecunicatedude propriosparamento sapiens Sinnia to ecunicatedude propriosparamento sapiens ATPase, aminophospholipid transporte Homo sapiens gene for AF-6, complete cds at 78h05.x1 Barstead aorta HPLRB6 Homo sapiens of Homo sapiens chromosome 21 unknown mRNA Homo sapiens chromosome 21 unknown mRNA Homo sapiens chromosome 21 unknown mRNA Homo sapiens plosphatase, target subunit 1 (H. sapiens gene encoding discoldin receptor tyrosine komo sapiens DNA for amyloid precursor protein, partial colluman prohormone converting enzyme (NEC2) gene, 801335244F1 NIH_MGC_39 Homo sapiens cDNA cit 601067378F1 NIH_MGC_10 Homo sapiens cDNA cit 601067378F1 NIH_MGC_10 Homo sapiens cDNA cit 601067378F1 NIH_MGC_10 Homo sapiens cDNA cit 601067378F1 NIH_MGC_10 Homo sapiens cDNA cit 601067378F1 NIH_MGC_29 Homo sapiens cDNA cit 601067378F1 NIH_MGC_20 Homo sapiens cDNA cit 601067378F1 NIH_MGC_29 Homo sapiens cDNA cit 601067378F1 NIH_MGC_20 Homo sapiens cDNA cit 601067378F1 NIH_MGC_20 Homo sapiens cDNA cit 601067378F1 NIH_MGC_20 Homo sapiens cDNA cit 601067378F1 NIH_MGC_20 Homo sapiens cDNA cit 601067378F1 NIH_MGC_20 Homo sapiens cDNA cit 601067378F1 NIH_MGC_20 Homo sapiens cDNA cit 601067378F1 NIH_MGC_20 Homo sapiens cDNA cit 601067378F1 NIH_MGC_20 Homo sapiens cDNA cit 601067378F1 NIH_MGC_20 Homo sapiens cDNA cit 601067378F1 NIH_MGC_20 Homo sapiens cDNA cit 601067378F1 NIH_MGC_20 Homo sapiens cDNA cit 601067378F1 NIH_MGC_20 Homo sapiens cDNA cit 601067378F1 NIH_MGC_20 Homo sapiens cDNA cit 601067378F1 NIH_MGC_20 Homo sapiens cDNA cit 601067378F1 NIH_MGC_20 Homo sapiens cDNA cit 601067378F1 NIH_MGC_20 Homo sapiens cDNA cit 601067378F1 NIH_MGC_20 Homo sapiens cDNA cit 601067378F1 NIH_MGC_20 Homo sapiens cDNA cit 601067378F1 NIH_MGC_20 Homo sapiens cdNA cit 601067378F1 NIH_MGC_20 Homo sapiens cdNA cit 601067378F1 NIH_MGC_20 Homo sapiens cdNA cit 601067378F1 NIH_MGC_20 Homo sapiens cdNA cit 601067378F1 NIH_MGC_20 Homo sapiens cdNA cit 601067378F1 NIH_MGC_20 Homo sapiens cdNA cit 601067378F1 NIH_MGC_20 Homo sapi	incophospholipid transition of complete cds -6, complete cds HPLRB6 Homo sapier e 21 unknown mRNA e 21 unknown mRNA sphatase, target subun iiscoldin receptor bytos yold precursor protein, part fring enzyme (NEC2); 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Table 4
Single Exon Probes Expressed in Heart.

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Top Hit Descriptor	AU118985 HEMBA1 Homo sapiens cDNA cione HEMBA1004795 61	Homo sapians myosin, heavy polypeptide 4, skeletal muscle (MYH4), mRNA	Homo sapiens amyloid beta (A4) precursor protein (protease nexin-II, Alzheimer disease) (APP), mRNA	Homo sapiens chromosome 21 unknown mRNA	Homo sapiens chromosome 21 unknown mRNA	Homo sapiens mRNA for T-box transcription factor (TBX20 gene), partial	Homo saplens mRNA for T-box transcription factor (TBX20 gene), partial	Homo sapiens ALR-like protein mRNA, partial cds	Homo saplens ALR-like protein mRNA, partial∣cds	Homo sapiens Kruppel-like factor 7 (ublquitous) (KLF7), mRNA	Homo saplens protein phosphatase 2A BR gamma subunit gene, exon 3	Homo saplens protein phosphatase 2A BR gaimma subunit gene, exon 3	601159563F2 NIH_MGC_53 Homo saplens cDNA clone IMAGE:3511118 5'	Homo sapiens similar to SALL1 (sal (Drosophila)-like (LOC57167), mRNA	Homo sapians chromosome 8 open reading frame 2 (C8ORF2), mRNA	Homo sapiens mRNA for KIAA0903 protein, partial cds	Homo saplens mRNA for KIAA0903 protein, partial cds	Homo sapiens soluble interfeukin 1 receptor accessory protein (IL1RAP) gene, exon 8, atternative exons 9 and complete ods. alternatively spliced	Homo sapiens mRNA for KIAA0633 protein, partial cds	Homo sapiens solute carrier family 1 (high affinity aspartate/glutamate transporter), member 6 (SLC1A6), mRNA	Homo sapiens brefeldin A-inhibited guanine nucleotide-exchange protein 2 (BIG2), mRNA	Homo sapiens SNCA isoform (SNCA) gene, complete cds, alternatively spliced	Homo sapiens CGI-15 protein (LOC51006), mRNA	Homo sapiens CGI-15 protein (LOC51006), mRNA	Homo sapiens DNA for Human P2XM, complete cds	Homo sapiens DNA for Human P2XM, complete cds	HUM000S381 Liver HepG2 cell line. Homo sapiens cDNA done s381 3'	Rattus norvegicus brain specific cortactin-binding protein CBP90 mRNA, partial cds	Homo sapiens makorin, ring finger protein, 1 (MKRN1), mRNA	zi90b04.s1 Soares_fetal_liver_spleen_1NFLS_iS1 Homo sapiens cDNA clone IMAGE:448015 3'	AU143539 Y79AA1 Homo saplens cDNA clone Y79AA1002087 5'
Top Hit Database Source	EST_HUMAN	FN	LΝ	NT	Ŋ	IN	IN	IN	1N	NT	IN	NT	EST_HUMAN	NT	N	NT	NT	IN	N	FZ	NT	NT	NT	LΝ	TN	NT	EST_HUMAN	NT	. LN	EST_HUMAN	EST_HUMAN
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Most Similar (Top) Hit BLAST E Value	2.0E-90	2.0E-90	1.0E-90	1.0E-90	1.0E-90	1.0E-90	1.0E-90	1.0E-90	1.0E-90	1.0E-90	1.0E-90	1.0E-90	1.0E-90	1.0E-90	1.0E-90	1.0E-90	1.0E-90	1.0E-90	1.0E-90	1.0E-90	1.0E-90	1.0E-90 A	1.0E-90	1.0E-90	1.0E-90	1.0E-90	8.0E-91	7.0E-91	7.0E-91	5.0E-91	5.0E-91
Expression Signal	1.56	49.27	3.39	1.21	1.04	2.03	2.03	1.71	7.71	2.45	3.47	3.47	4.02	4.98	8.4		1.18	1	2.2	2.85	3.78	1.22	1.72	1.72	1.89	1.89	5.48	0.88	2.05	1.47	1.05
ORF SEQ ID NO:	27797	28159	20060	20148	20148	20435	20438	20472	20473					21631	22645			24005		26558			27444	27445		26226		21189	26907		24092
Exan SEQ ID NO:	17572	17914	10241	12639	12639	10613	10613	10645	10645	11010	11192	11192	11548	11756	12752	13689	13689	14223		16380	16998	17224	17240	17240		19268	14001	11333	16714		14309
Probe SEQ ID NO:	77.22	8765	275	370	371	089	089	713	713	1094	1284	1284	1644	1860	2823	3777	3777	4326	5481	6521	7121	7356	7371	7371	9732	9732	4101	1428	6835	3429	4415

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Table 4
Single Exon Probes Expressed in Heart

		Т	Г	Г	Г	П		П			П										m	<u> </u>	Π		T			Ė	T		
	Top Hit Descriptor	AU143539 Y78AA1 Homo sapiens cDNA clone Y79AA1002087 5'	Homo sapiens chromosome 22 open reading frame 5 (C22ORF5), mRNA	Homo sapiens chromosome 22 open reading/frame 5 (C22ORF5), mRNA	AV649878 GLC Homo sapiens cDNA clone (3LCBYF08 3'	AV649878 GLC Homo sapiens cDNA clone GLCBYF08 3'	Homo saplens lysophosphatidic acid acyltransferase-delta (LPAAT-delta) mRNA, complete cds	Homo sapiens lysophosphatidic acid acytranisferase-delta (LPAAT-delta) mRNA, complete cds	Homo saplens chromosome 21 segment HS21C084	EST01579 Hippocampus, Stratagene (cat. #936205) Homo sapiens cDNA clone HHCMC60 similar to Retrovirus-related gag polyprotein	EST01579 Hippocampus, Stratagene (cat. #836205) Homo sapiens cDNA clone HHCMC60 similar to	Retrovirus-related gag polyprotein	Homo sapiens solute carrier family 4, anion exchanger, member 3 (SLC4A3), mRNA	Homo sapiens solute carrier family 4, anion exchanger, member 3 (SLC4A3), mRNA	Homo sapiens beta-ureldopropionase (BUP1) gene, exon 6	Homo sapiens beta-ureidopropionase (BUP1) gene, exon 6	Homo sapiens chromosome 21 segment HS2/1C083	Homo saplens mRNA for KIAA1278 protein, partial cds	Homo sapiens mRNA for KIAA1278 protein, partial cds	Homo sapiens cyclin-D binding Myb-like protein mRNA, complete cds	Human Ku (p70/p80) subunit mRNA, complete cds	Homo sapiens chromosome 21 segment HS2/C085	Homo capiens chromosome 21 segment HS2/i C085	Homo sapiens epididymal secretory protein (19.5kD) (HE1), mRNA	Hamo sapiens cyclin-dependent kinase 6 (CDK6) mRNA	Homo sapiens gamma-aminobutyric acid (GAI3A) B receptor, 1 (GABBR1), transcript variant 2, mRNA	Homo sapiens gamma-aminobutyric acid (GABA) B receptor, 1 (GABBR1), transcript variant 2, mRNA	Human L-type calcium channel beta-1 subunit (CACNLB1) gene, exons 10 and 11	Human L-type calcium channel beta-1 subunit (CACNLB1) gene, exons 10 and 11	Human mRNA for very low density lipoprotein receptor, complete cds	Homo sepiens glutathione S-transferase theta?2 (GSTT2) and glutathione S-transferase theta 1 (GSTT1) genes, complete cds
	Top Hit Database Source	EST_HUMAN	LΝ	Į.	EST_HUMAN	EST_HUMAN	M	FZ	N	EST HUMAN		EST_HUMAN	Ν	NT	LN	N	IN	TN	NT	NT	NT	NT	NT	NT	NT	TN	IN	NT	NT	NT	NT
-	Top Hit Acesslan No.	AU143539.1	7110634 NT	7110634 NT	AV649878.1	AV649878.1	AF156776.1	AF156776.1	AL163284.2	M77994.1		M77894.1	114301B3 NT	11430193 NT	AF169555.1	AF169555.1	AL163283.2	AB033104.1	AB033104.1	AF084530.1	M30938.1	AL163285.2	AL163285.2	11434964 NT	4502740 NT	11497611 NT	11497611 NT	U86959.1	U86959.1	D16494.1	AF240786.1
	Most Similar (Top) Hit BLAST E Value	5.0E-91	_	5.0E-91	_		4.0E-91	4.0E-91	4.0E-91	4.0E-91		4.0E-91	3.05-91	3.0E-91				3.0E-91	3.0E-91		3.0E-91	3.0E-91		3.0E-91	3.0E-91	3.05-91	3.0E-91		3.0E-91		3.0E-91
	Expression Signal	1.05	0.82	0.82	1.34	1.34	1.3	1.3	3.13	1.67		1.67	5.12	5.12	68.0	0.99	1.77	2.96	2.96	0.83	4.02	1.2	1.2	1.45	2.39	4.11	4.11	4.4	4.4	3.31	1.45
	ORF SEQ ID NO:	24083	24379				22895	22896	28427	25314		25360		21367	22382			23136	23137		24168	24560	24561	25470		25996	25997	L			25286
	Exan SEQ ID NO:	14309	14589	14589	16964	16964	13091	13091	18180	18954	ı	18954	11506	11506	12492	12492	13219	13333	13333		14381	14785	14785	15407	15712	15872	15872			16970	19104
	Probe SEQ ID NO:	4415	4703	4703	7087	7087	3166	3166	8301	9239		9239	1601	1691	2624	2624	3297	3416	3416	3720	4487	4905	4905	5488	2807	5967	2962	6502	6502	7083	8480

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Hamo sapiens dihydrolipoamide S-succinyltransferase (E2 component of 2-oxo-glutarate complex) (DLST) Homo sapiens nuclear mitogen- and stress-activated protein kinase-1 (MSK1) mRNA, complete cds UI-H-BI3-aks-d-01-0-UI.s1 NCI_CGAP_Subs Homo sepiens cDNA clone IMAGE:2735280 Homo sapiens brefeldin A-Inhibited guanine nucleotide-exchange protein 2 (BIG2), mRNA ym30e03.r1 Scares Infant brain 1NIB Homo sapiens cDNA clone IMAGE:49587 5' 602022088F1 NCI_CGAP_Brn67 Homo sapiens cDNA clone IMAGE:4157804 5' 602022088F1 NCI_CGAP_Brn67 Homo sapiens cDNA clone IMAGE:4157804 5' Homo sapiene fragile X mental retardation, autosomal homolog 1 (FXR1), mRNA Human von Willebrand factor pseudogene corresponding to exons 23 through 34 Homo saplens mRNA for KIAA0758 protein, partial cds Homo saplens cytoplasmic Seprase truncated isoform mRNA, complete cds 801273513F1 NIH_MGC_20 Homo saplens cDNA clone IMAGE:3614687 6 26f3 Human retina cDNA randomly primed sublibrary Homo sapiens cDNA Homo sapiens diacy/glycerol kinase, gamma (90kD) (DGKG), mRNA Homo sapiens diacy/glycerol kinase, gamma (90kD) (DGKG), mRNA Homo saplens hypothetical protein FLJ20260 (FLJ20260), mRNA Top Hit Descriptor 10mo sapiens beta-ureidopropionase (BUP1) gene, exon 6 Homo sapiens beta-ureidopropionase (BUP1) gene, exon 6 Human Na+,K+ ATPase alpha-subunit mRNA, partial cds Homo sapiens B-cell CLL/lymphoma 7b (BCL7B) mRNA Homo sapiens mRNA for KIAA0758 protein, pertial ods Homo sapiens mRNA for KIAA1512 protein, partial cds Homo sapiens mRNA for KIAA0611 protein, partial cds Homo sapiens mRNA for MBNL protein Homo sapiens mRNA for KIAA1512 protein, partial ods Human lens membrane protein (mp19) gene, exon 11 Homo sapiens chromosome 21 segment HS21C084 Human lens membrane protein (mp19) gene, exon Homo sapiens NALP1 mRNA, complete cds Homo sapiens NKG2D gene, exon 10 Homo sapiens NKG2D gene, exon 10 Homo sapiens MCP-4 gene mRNA EST_HUMAN HUMAN EST_HUMAN EST HUMAN EST_HUMAN HUMAN Top Hit Database Source 눋 되보토 눋 눌 4503340|NT 눋 11434722|NT 占 눌 4502384 NT 11434704 11422086 11427149 11434722 Top Hit Acession 9.0E-92 AB040945.1 9.0E-92 AB040945.1 7.0E-92 AB018301.1 AL163284.2 AW 449746.1 9.0E-92 AF310105.1 8.0E-92 AF074393.1 BF348182.1 8.0E-92|BE386363.1 AJ000979.1 AF169555.1 7.0E-92 AF007822.7 7.0E-92 46 9.0E-92 AJ001689.1 1.0E-91 H15212.1 9.0E-92 AJ001689.1 AB014511. ģ 7.0E-92 M60676.1 Y13829.1 8.0E-92 L04193.1 8.0E-92 L04193.1 8.0E-92 3.0E-91 3.0E-91 1.0E-91 8.0E-92 8.0E-92 8.0E-92 1.0E-91 1.0E-91 8.0E-92 8.0E-92 (Top) Hit BLAST E Aost Similar Value -2.51 2.64 3.22 19.33 53. 54. 2.91 1.91 8.41 19.33 87 3.61 3.2 3.02 8.41 Expression Signal 28318 19858 20020 26176 25268 21017 ORF SEQ ID NO: 19833 20987 26176 25418 25911 27409 20088 21554 21555 26978 20021 20981 19887 27836 18556 12662 12662 11166 16035 16035 10248 11677 16786 16786 17610 18067 19162 1889 15362 15789 10071 11677 SEO ID 18757 16757 ÿ 1259 5882 6878 1778 9812 9812 5353 8289 8069 8180 9572 236 238 3 9398 22 5442 87 283 5944 6908 8667 80 SEQ ID Probe ë

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	_	_				_	_				_				-				-			*	II. Weller	7917			*****		P "MES" (AMI
Top Hit Descriptor	Homo sapiens ARP2 (actin-related protein 2, yeast) homolog (ACTR2), mRNA	Homo sapiens ARP2 (actin-related protein 2, yeast) homolog (ACTR2), mRNA	Homo saplens cysteine-rich repeat-containing protein S52 precursor, mRNA, complete cds	Homo sapiens NRAS-related gene (D1S155E), mRNA	Homo sapiens DNA, MHC class I region, 7.1 encestral haplotype	Homo sapiens T-cell lymphoma invasion and metastasis 1 (TIAM1) mRNA	Homo sapiens T-cell lymphoma invasion and metastasis 1 (TIAM1) mRNA	N-CAM=145 kda neural cell adhesion molecule [human, small cell lung cancer cell line OS2-R, mRNA, 2980 nt]	N-CAM=145 kda neural cell adhesion molecule [human, small cell lung cancer cell line OS2-R, mRNA, 2960 nt]	Homo saplens prospero-related homeobox 1 (PROX1) mRNA	zw66d12.r1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:781175 5'	601283012F1 NIH_MGC_44 Homo sapiens ci2NA clone IMAGE:3605018 6'	601501242F1 NIH_MGC_70 Homo sapiens cDNA clone IMAGE:3902939 5'	EST91020 Synovial sarcoma Homo sapiens cl0NA 5' end similar to similar to ribosomal protein S13	Human mRNA for alpha-actinin	Human mRNA for alpha-actinin	Homo sapiens activin A receptor, type IIB (ACVR2B) mRNA	Homo sapiens hypothetical protein dJ462O23.2 (DJ462O23.2), mRNA	Homo saplens hypothetical protein dJ462O23.2 (DJ462O23.2), mRNA	601118337F1 NIH_MGC_17 Homo sapiens c0NA clone IMAGE:3028304 5	601118337F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:3028304 6	mrg=mas-related [human, Genomic, 2416 nt]	wk27d07.x1 NCI_CGAP_Bm25 Homo sapiens cDNA clone IMAGE:2413549 3' similar to TR:Q12844 O12844 BREAKPOINT CLUSTER REGION PROTEIN	W.Z7407.x1 NCI_CGAP_Brn.25 Homo saplens cDNA clone IMAGE:2413549 3' similar to TR:Q12844 Q12844 BREAKPOINT CLUSTER REGION PROTEIN :	Homo sapiens syndecan 4 (amphiglycan, ryudocan) (SDC4) mRNA	Homo sapiens calcineurin binding protein 1 (KiAA0330), mRNA	Homo sapiens chromosome 21 unknown mRNA	Homo sapiens chromosome 21 unknown mRNA	Homo sapiens stress-induced-phosphoprotein 1 (Hsp70/Hsp90-organizing protein) (STIP1), mRNA
Top Hit Database Source	NT	ĮN	NT	N	NT	FZ	LN	Į.	L	ΝŢ	EST HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	2	N	LN	ΙN	NT	EST_HUMAN	EST_HUMAN	LN LN	EST HUMAN	EST HUMAN	TN	L'N	TN	TN	NT
Top Hit Acession No.	5031570 NT	5031570 NT	AF167706.1	E006738 NT	AB031007.1	4507500 NT	4507500 NT	S71824.1	S71824.1	4508118	AA446208.1			AA378336.1	X15804.1	X15804.1	4501898 NT	11422946 NT	11422946 NT	BE299190.1	BE299190.1	S78653.1	A1818119.1	Al818119.1	09890	6912457 NT	AF231919.1	AF231919.1	58031B0 NT
Most Similar (Top) Hit BLAST E Value	7.0E-92	7.0E-92	7.0E-92	7.0E-92	7.0E-92	7.0E-92	7.0E-92	7.0E-92			7.0E-92	6.0E-92	3.0E-92	3.0E-92	3.0E-92	3.0E-92)	2.0E-92	2.0E-92	2.0E-92	2.0E-92	2.0E-92	2.0E-92	2.0E-92	2.0E-92	2.0E-92	2.0E-92	2.0E-92	2.0E-92	2.0E-92
Expression Signal	1.67	1.67	2.32	5.01	1.04	0.92	0.92	1.61	1.61	1.45	4.87	0.83	2.45	3.74	5.7	5.7	1.53	2.93	2.93	1.38	1.38	2.74	1.55	1.55	4.71	37.64	1.02	1.02	4.99
ORF SEQ ID NO:	Ц	21924	22283	22445	22481	23023	23024	24165	24166			L		25575			19803	19958	19959		20498		21667	21668	21787			23266	23332
Exen SEQ ID NO:	1 1	12027	12391	12558	12586	15068	15068	14378	14378		L	L	L	L	18034	18034	10010	10143	10143	10664	10664	11588	11789	11789	11895		L	13476	13545
Probe SEQ ID NO:	2139	2139	2517	2683	2724	3301	3301	4484	4 84	5110	5219	1569	2738	5583	8146	8146	23	172	172	732	732	1686	1894	1894	2002	2623	3562	3562	3831

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Top Hit Descriptor	Human endogenous retroviral DNA (4-1), complete retroviral segment	Homo sapiens bile salt export pump (BSEP) mRNA, complete cds	UNFZP444U414 11 434 (synonym: ntess) Homo sapiens cUNA clane UNFZP434U0414 5	Homo sapiens mRNA for KIAA1068 protein, partial cds	Human NPY Y1-like receptor pseudogene mRNA, complete cds	hd02h02.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2908371 3' similar to TR:O02711 002711 PRO-POL-DUTPASE POLYPROTEIN ;	Homo sapiens thyroid stimulating hormone receptor (TSHR), mRNA	Homo sapiens mRNA for KiAA1093 protein, partial cds	Homo sapiens calcineurin binding protein 1 (KIAA0330), mRNA	y/80e08.r1 Soares placenta Nb2HP Hamo sapiens cDNA clone IMAGE:145574 5'	y80e08.r1 Scares placenta Nb2HP Homo sapiens cDNA clone IMAGE:145574 5'	Homo sapiens ribosomal protein, large, P1 (RPLP1) mRNA	tg01b02.x1 NCI_CGAP_CLL1 Home sapiens cDNA clone IMAGE.2107487 3' similar to SW.PTNF_HUMAN Q16825 PROTEIN-TYROSINE PHOSPHATASE D1 ; contains Alu repetitive element; contains element	MER17 repetitive element;	tg01b02.x1 NCI_CGAP_CLL1 Home sapiens CDNA clone IMAGE:2107467 3' similar to SW:PTNF_HUMAN Q16825 PROTEIN-TYROSINE PHOSPHATASE D1; contains Alu repetitive element; contains element MED17 repetitive element.	אוריוון בליפחוא ממונפון	AU121681 MAMMA1 Homo capiens cDNA cicne MAMMA1000738 5	EST188414 HCC cell line (matastasis to liver in mouse) II Homo sapiens cDNA 5' end similar to ribosomal protein L29	Homo sapiens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively	spliced	601281867F1 NIH_MGC_44 Homo saplens cDNA clone IMAGE:3603832 5'	Homo sapiens ribosomal protein L10a (RPL10A), mRNA	601460521F1 NIH_MGC_66 Homo saplens oldNA clone IMAGE:3863908 5'	Homo saplens chromosome 21 unknown mRNA	Homo sapiens mRNA for KIAA0611 protein, partial cds	wc09c08.x1 NCI_CGAP_Pr28 Homo sapiens cDNA clone IMAGE:2314670 3'	wc09c08.x1 NCI_CGAP_Pr28 Homo sapiens cDNA clone IMAGE:2314670 3'	Homo sapiens chromosome 21 segment HS21C001	Human skeletal muscle 1.3 kb mRNA for tropomyosin
Top Hit Database Source	닐	- L	ESI HOMAN	Ł	L Z	EST_HUMAN	Z	FN	LN LN	EST_HUMAN	EST_HUMAN	N		EST_HUMAN	NAME IN TOO	בסו בטולואוט	EST HUMAN	EST_HUMAN		NT	EST_HUMAN	NT	EST_HUMAN	FX	NŦ	EST_HUMAN	EST_HUMAN	N-I	NT
Top Hit Acession No.	M10976.1	AF136523.1	ALU4043/.1		U67780.1	AW340174.1	11434900 NT	AB029016.1	6912457 NT	R78078.1	R78078.1	450668 NT		AI380358.1	. SHOUDGIV	Alsowood. 1	AU121681.1	AA316723.1		AF223391.1	BE388571.1	11418526 NT	BF036364.1	AF231919.1	AB014511.1	AI674184.1	AI674184.1	AL163201.2	X04201.1
Most Similar (Top) Hit BLAST E Value	2.0E-92	2.0E-92	Z0E-92	2.0E-92	2.0E-92	2.0E-92	2.0E-92	2.0E-92	2.0E-92	1.0E-92	1.0E-92	1.0E-92		1.0E-92	20 10		9.0E-93	9.0E-93		9.0E-93	9.0E-93	9.0E-93	8.0E-93	7.0E-93	5.0E-93	5.0E-93	5.0E-93		5.0E-93
Expression Signal	1.02	0.79	7.33	2.49	2.25	1.37	5.91	2.55	26.65	1.11	1.11	34.72	· • · • •	4.04	70 7	5	3.14	9.21		1.46	96.0	8.8	2.49	6.24	1.25	62.39	5.39	0.95	2.42
ORF SEQ ID NO:		24323		26017		27211	28276		22381	21582	21583			27360	19646	١	21769				23267		26003	20028	21121	21146	21148		22925
Exan SEQ ID NO:			_[16273	17018	18030	19172	12491	11704	11704	11919		17162	47482	ı	11876	11890		12478	13477	18719	15879	10212	11265	11290	11290		13120
Probe SEQ ID NO:	4191	4648	4322	5983	9420	7141	8142	9589	6836	1807	1807	2028		7286	7786	207/	1983	1896		2610	3563	8911	9269	246	1359	1385	1385	1459	3195

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	Top Hit Descriptor	Homo sapiens tensin mRNA, complete cds	Human Olk-associated RS cyclophilin CARS-Gyp mRNA, complete cds	601117586F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3358220 5	601116810F1 NIH_MGC_16 Homo sapiens ciDNA clane IMAGE:3357243 5'	EST376458 MAGE resequences, MAGH Homo saplens cDNA	Homo sapiens hypothetical protein (LOC51318), mRNA	UI-HF-BND-aks-g-09-0-UI.r1 NIH_MGC_50 Homo sapiens cDNA clone IMAGE:3078329 5'	qp78b10.x1 Soares_fetal_lung_NbHL19W Homo saplens cDNA clone IMAGE:1929115 3'	229c10.s1 Soares_pregnant_uterus_NbHPU Homo sapiens cDNA clone IMAGE:503346 3'	Homo sapiens CYP17 gene, 5' end	601458531F1 NIH_MGC_66 Homo sapiens cDNA clane IMAGE:3862086 6	Homo sapiens CTR1 pseudogene	Homo sapiens CTR1 pseudogene	Homo sapiens hypothetical protein (DJ328E19, C1.1), mRNA	oy84b08.x1 NCI_CGAP_CLL1 Homo saplens cDNA clone IMAGE:1672503 3' similar to TR:Q62384 Q62384	Homo sanjana DNA for amyloid macursor profision complete cole	Home captains broadhaited profess Project on Participation	Hallo Sapieris Hypotrieucal progent Lassace (1 Casace), minto	Homo saplens hypothetical protein FL/20291 (FL/20291), mRNA	Homo sapiens mRNA for KIAA1563 protein, partial cds	Homo saplens cysteine-rich repeat-containing protein S52 precursor, mRNA, complete cds	Homo sablens long chain polyunsaturated fatty acid elongation enzyme (HELO1) mRNA, complete cds	Homo saplens MHC class 1 region	Novel human gene mapping to chomosome 1	601177686F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:3532965 5'	601177686F1 NIH_MGC_17 Homo sapiens d0NA clone IMAGE:3532965 5'	Homo sapiens DNA for amyloid precursor protein, complete cds	Homo sapiens chromosome 21 segment HS21C084	Homo sapiens glucocorticoid receptor (GRL) gene, intran D, exon 5, and intron E	Homo sapiens glucocorticoid receptor (GRL) gene, intron D, exon 5, and intron E	Homo sapiens neurofibromin 1 (neurofibromatosis, von Recklinghausen disease, Watson disease) (NF1) mRNA	Homo sapiens protein kinase C, beta 1 (PRKCB1), mRNA
1 5	Database Source	۲ ۲	ΓN	EST_HUMAN	EST_HUMAN	EST_HUMAN	LZ LZ	EST_HUMAN	EST_HUMAN	EST_HUMAN	Į.	EST_HUMAN	NT	LΝ	TN	MANUEL FOR	LSI DONAN	1	2	LV.	NT	NT	ΙΛ	Ę	L	EST_HUMAN	EST HUMAN		NT	NT.	ΝT	Ę	NT
	Top Hit Acession No.	AF225896.1	U40763.1	BE252982.1	BE253201.1	AW964385.1	11430039 NT	AW 502002.1	Al312025.1	AA126735.1	L41825.1	BF035327.1	AF238997.1	AF238997.1	7657016 NT	NIA ACTOE A	A1140733.1	TIMIOTOCOG	0923270	8923270 NT	AB046783.1	AF167706.1	AF231981.1	AF055066.1	AL137200.1	BE297369.1	BE297369.1	D87675.1	AL163284.2	U78509.1	U78509.1	4657792 NT	11431590 NT
Most Similar	(Top) Hit BLAST E Value		2.0E-93	2.0E-93	2.0E-93	2.0E-93 /	2.0E-93	2.0E-93 /	2.0E-93	2.0E-93	2.0E-93		1.0E-93	1.0E-93	1.0E-93		1.0F-93		1.05-93		1.0E-93 /	1.0E-93 /	1.0E-93		1.0E-93	1.0E-93	1.0E-93	1.0E-93	1.0E-93	1.0E-93	1.0E-93	1.0E-93	1.0E-93
	Expression Signal	1.48	1.02	0.89	1.02	4.59	1.52	1.32	2.87	1.52	1.31	2.75	1.64	1.64	2.58	27.0	3.73	47.0	0.41	6.41	1.65	1.68	1.01	4.16	1.09	2.69	2.69	4.33	1.44	1.62	1.62	9.15	2.08
	ORF SEQ ID NO:		21868	22210	24732	25049	25442		29093				19897	19898	20282		20834					21085		22190		21030	21031		24010	26391	25392	26604	26197
100	<i>o</i>	11503	11973	12313	L	15245	15382		19476	19046	19095	19263	10081	10081	10449		10523	L		11125	11228	11230	12172	L		11181	11181	12830	14228	15338	15338	16522	16051
Drohe	SEQ ID NO:	1598	2083	2436	5088	5325	5462	6014	9668	9388	9465	9724	96	96	209	100	282 854	1574	2	1217	1321	1323	2289	2415	2459	2782	2782	2903	4331	5417	5417	5607	6067

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	Top Hit Descriptor	Human mRNA for NF1 N-isoform-exon11, complete cds	Homo saplens mRNA for KIAA1411 protein, partial cds	H.capiens mRNA for MEMD protein	Homo sapiens protein kinase inhibitor gamma (PKIG) mRNA, complete cds	Homo saplens mRNA for KIAA1485 protein, pertial cds	Homo sapiens Trio isoform mRNA, complete cds	Human PreA4 gene for Alzheimer's disease A4 amyloid protein precursor (exon 9)	Human PreA4 gene for Alzhelmer's disease A4 amyloid protein precursor (exon 9)	qm03c12x1 NCI_CGAP_Lu5 Hamo sapiens cDNA clone IMAGE:1880758 3' similar to WP:T19B4.4 (CE13742 ;	Homo saplens glutathlone S-transferase theta 2 (GSTT2), mRNA	Homo sapiens chromosome 21 segment HS21C009	Homo sapiens transcription enhancer factor-5 inRNA, complete cds	Homo saplens mitogen-activated protein kinase 12 (MAPK12), mRNA	Homo sapiens mRNA for KIAA0612 protein, partal cds	Homo sapiens mRNA for KIAA0612 protein, partial cds	zg87g06.s1 Soares_fetal_heart_NbHH19W Homo sapiens cDNA clone IMAGE:409594 3'	ot83d05.s1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA clone IMAGE:1623359 3'	yd98b04.s1 Soares fetal liver spleen 1NFLS Home sapiens cDNA clone IMAGE:116239 3'	Homo saplens mRNA for KIAA0027 protein, partial cds	Homo sapiens cleavage and polyadenylation specific factor 1, 160kD subunit (CPSF1), mRNA	Homo sapiens ribosomal protein L27 mRNA, complete cds	Homo sapiens protein phosphatase 1, regulatory subunit 10 (PPP1R10) mRNA	tw11f10.x1 NCI_CGAP_Brn52 Hamo septens cDNA clone IMAGE:2259403 3' similar to TR:Q15265 Q15265 PROTEIN TYROSINE PHOSPHATASE;	Homo sapiens solute carrier family 22 (organic cation transporter), member 1-like (SLC22A1L), mRNA	Homo sapiens solute carrier family 22 (organic cation transporter), member 1-like (SLC22A1L), mRNA	Homo saplens hypothetical protein FLJ12455 (FLJ12455), mRNA	Homo sapiens ASH2L gene, complete cds, striilar to Drosophila ash2 gene	Homo sapiens complement component 6 (C5)/mRNA	Homo saplens cystelne-rich repeat-containing protein S52 precursor, mRNA, complete cds	Homo sapiens cysteine-rich repeat-containing protein S62 precursor, mRNA, complete cds
,	Top Hit Database Source	TN	NT	TN	LN	NT	NT	TN	IN	EST HUMAN	N-	LN LN	Z	之	LN	IN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	TA	ΝΤ	NT	EST_HUMAN	Z	Ę	<u> </u>	Z	TN	Z	뉟
	Tap Hit Acession No.	D42072.1	AB037832.1	Y10183.1	AF182032.1	AB040918.1	AF091395.1	X13474.1	X13474.1	Al268262.1	11417856 NT	AL163209.2	AF142482.1	11418351 NT	AB014512.1	AB014512.1	AA722434.1	AI015800.1	T89398.1	D25217.2	9558724 NT	L05094.1	4506008 NT	AI591312.1	11440670 NT	11440870 NT	11545792 NT	AB022785.1	4502508 NT	AF167706.1	AF167708.1
	Most Similar (Top) Hit BLAST E Value	1.0E-93	1.0E-93	1.0E-93	1.0E-93	1.0E-93 /		1.0E-93	1.0E-93.7	1.0E-93				6.0E-94	5.0E-94	5.0E-94	5.0E-94	5.0E-94	5.0E-94	5.0E-94	5.0E-94	4.0E-94	4.0E-94	4.0E-84	4.0E-94	4.0E-94	4.0E-94	3.0E-94	3.0E-94		3.0E-94
	Expression Signal	4.09	2.04	1.18	1.59	1.8	1.22	4:54	4.64	5.92	2.33	122	1.74	1.67	3.05	3.05	1.72	1.63	4.68	1.27	1.26	4.55	98.0	3.02	1.84	1.84	1.72	1.17	1.17	1.05	1.05
	ORF SEQ ID NO:	26320			L	26640		27589		25132			23579		25019	25020		26218	24911				22379	24295	25921					Ц	21478
	Exan SEQ ID NO:	16163	1_	16848	16901	16450	16453	17380	17380				L.		15217	15217	15598	16069	19736	19378	19384	11697	12489	14508	15798	I		L		1706 11607	11607
	Probe SEQ ID NO:	6588	6807	1.769	7024	7437	7440	7529	7529	9051	9716	8026	3880	9830	5298	5296	5689	6183	9361	9901	9907	1799	2621	4618	5892	5892	8749	589	704	1706	1706

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Table 4
Single Exon Probes Expressed in Heart

1		ī	Т	1	Т	т	Т	Т	1	Т	Т	Т	Т	П	Т	1		T.	Ť ~	Ť	T -	i	T	1	Т	T	_	т-	_	Ė	Ė
ongo the second religion	Top Hit Descriptor	Homo saplens E1A binding protein p300 (EP300) mRNA	zw63g08.r1 Soares_total_fetus_Nb2HF8_9w Home sapiens cDNA clone IMAGE:774782 5'	Homo saplens zinc finger protein 277 (ZNF277), mRNA	Homo saplens chromosome 21 open reading frame 18 (C21ORF18), mRNA	Homo sapiens protocadherln alpha 13 (PCDH-alpha13) mRNA, complete cds	Homo sapiens mRNA for KIAA0879 protein, partial cds	Homo sapiens glycogenin-1L mRNA, complete cds	Homo sapiens axonal transport of synaptic vesicles (ATSV) mRNA	Human cbl-b truncated form 1 lacking teucine zipper mRNA, complete cds	601175762F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:3531038 5'	601111698F1 NIH_MGC_16 Homo saplens ciONA clone IMAGE:3352559 5'	601111696F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3352559 5	Homo sapiens hypothetical protein (FLJ20746), mRNA	Homo sapiens paired box gene 5 (B-cell lineage specific activator protein) (PAX5), mRNA	601468748F1 NIH_MGC_67 Homo sapiens cIDNA clone IMAGE:3872099 5'	Homo sapiens IL-1 receptor antagonist IL-1Raj(IL-1RN) gene, alternatively spiloed forms, complete cds	apz2e02.xt Schiller oligodendroglioma Homo capiens cDNA clone IMAGE:1956122 3' similar to TR:062945 062845 NEURAL CELL ADHESION PROTEIN BIG-2 PRECURSOR	601175762F1 NIH MGC_17 Homo saplens cDNA clone IMAGE:3531038 5'	Homo sapiens TNF-alpha stimulated ABC protein (ABC50) mRNA, complete cds	Homo sapiens KIAA0255 gene product (KIAA0255), mRNA	Homo sapians KIAA0255 gene product (KIAA0255), mRNA	Homo sapians progressive ankylosis-like protein (ANK) mRNA, complete cds	we09e04x1 NCI_CGAP_Lu24 Homo saplens :DNA done IMAGE;2340606 3' similar to gb:K00558 TUBULIN ALPHA-1 CHAIN (HUMAN);	we09e04x1 NCI_CGAP_Lu24 Homo saptens :DNA clone IMAGE:2340606 3' similar to gb:K00558 TUBULIN ALPHA-1 CHAIN (HUMAN);	Homo saplens proteasome (prosome, macropain) 26S subunit, non-ATPase, 11 (PSMD11), mRNA	Homo saplens proteasome (prosome, macropain) 26S subunit, non-ATPase, 11 (PSMD11), mRNA	Homo saplens potassium channel subunit (HERG-3) mRNA, complete cds	Homo sapiens KIAA0255 gene product (KIAA0255), mRNA	Homo saplens KIAA0255 gene product (KIAA0255), mRNA	Homo sapiens proline dehydrogenase (proline oxidase) (PRODH) mRNA
51 1104 55	Top Hit Database Source	뉟	EST_HUMAN	Ā	ZT	LN LN	LN LN	١	۲N	TN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	N	EST_HUMAN	ΙN	EST HUMAN	EST HUMAN	FZ	NT	NT	TN	EST_HUMAN	EST HUMAN	NT	NT	LZ.	NT	NT	NT
	Top Hit Acession No.	4557556 NT	AA464805.1	11496268 NT	11526228 NT	AF152309.1	AB014579.1	AF087942.1	4757821 NT	U26711.1	BE295714.1	BE253433.1	BE253433.1	9506682	11428710 NT	BE780478.1	U65590.1	AI272244 1	BE295714.1	AF027302.1	7662027	7662027 NT	AF274763.1	AI700998.1	AI700998.1	11426529 NT	11426529 NT	AF032897.1	11420944 NT	11420944 NT	5174644 NT
	Most Similar (Top) Hit BLAST E Value	3.0E-94	3.0E-94	3.0E-94	3.0E-94		3.0E-94		3.0E-94	3.0E-94 L	1.0E-94		1.0E-94	1.0E-94	1.0E-94	1.0E-94	1.0E-94 L	1.0E-94	_		9.0E-95	9.0E-95	9.0E-95 /	8.0E-95	8.0E-95		8.0E-95		8.0E-95	8.0E-95	8.0E-95
	Expression Signal	3.45	0.82	3.58	4.16	1.18	3.79	4.36	1.75	2.27	2.24	1.91	1.91	1.13	1.83	1.41	2.49	2.19	1.98	1.55	1.13	1.13	1.87	1.59	1.59	1.83	1.83	2.05	1.73	1.73	2.82
	ORF SEQ ID NO:	21604	23772	25466	25909	26838	27062	27698	28580			22769	22770		27439		28546	28775	L				26869	24119	24120			26837			27744
	Exan SEQ ID NO:		13995	15403	15787	16650	16869	17384							17235	17487	18292	18500	L	11366	13043	13043	16680	14331	14331	16158		16649	Ц		17517
	Probe SEQ ID NO:	1735	4095	5484	5881	6771	6992	7633	8448	8928	143	3050	3050	4261	7331	7636	8418	8635	9759	1461	3118	3118	6801	4436	4436	6294	6294	6770	7391	7391	7667

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. Top Hit Descriptor	Homo sapiens mRNA for KIAA1395 protein, pariial cds	Homo saplens developmental arteries and neural crest EGF-like protein mRNA, complete cds	2u84b01.s1 Soares_testis_NHT Homo saplens cDNA clone IMAGE:744649 3' similar to contains L1.t1 L1 repetitive element:	Homo sapiens DNA for amyloid precursor protein, complete cds	Homo saplens DNA for amyloid precursor protein, completa cds	Homo saplens Ly-6-like protein (CD59) mRNA! complete cds	Homo saplens chromosome 21 segment HS21C046	Human homeobox protein (PHOX1) mRNA, 3' end	602071146F1 NCI_CGAP_Brn64 Homo sapiens cDNA clone IMAGE:4214147 5'	Homo sapiens H factor 1 (complement) (HF1) mRNA	Homo saplens KiAA0255 gene product (KiAA0255), mRNA	Homo saplans KIAA0255 gene product (KIAA0255), mRNA	Homo sapiens tissue inhibitor of metalloproteinase 3 (Sorsby fundus dystrophy, pseudoinflammatory) (TIMP3)	UNITED TO SERVICE STATE OF THE	601312161F1 NIH_MGC_44 Homo sapiens clONA clone IMAGE:3658862 5'	Homo sapiens G protein-coupled receptor 19 (SPR19) mRNA	Homo saplens G protein-coupled receptor 19 (GPR19) mRNA	Homo saplens glutathione S-transferase theta 2 (GSTT2) and glutathlone S-transferase theta 1 (GSTT1) genes, complete cds	Homo saplans glycha cleavage system protein H (aminomethy/ carrier) (GCSH) mRNA	Homo saplens H factor 1 (complement) (HF1) mRNA	Homo sapiens Usurpin-gamma mRNA, complete cds	Homo sapiens unconventional myosin-15 (LOC51168), mRNA	Homo sapiens unconventional myosin-15 (LOC51168), mRNA	Homo sapiens mRNA for KIAA1386 protein, partial cds	qm01c02x1 Soares_NhHMPu_S1 Homo capitans cDNA clone IMAGE:1880646 3' almilar to WP:T23G7.4 CE03706;	Homo saplens hypothetical protein (HS322B1A), mRNA	Homo saplens KIAA0187 gene product (KIAA0187), mRNA	zx11d07.r1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA clone IMAGE:786157 5/	zx11d07.r1 Soares_total_fetus_Nb2HF8_9w Homo saplens cDNA clone IMAGE:786157 5	Homo saplens CGI-48 protein (LOC51096), mRNA	Homo saplens CGI-48 protein (LOC51096), mRNA
Top Hit Database Source	LN L	NT		NT	NT	L	N L	LN L	EST_HUMAN						HUMAN			NT						IN	EST HUMAN			EST_HUMAN	EST_HUMAN		
Top Hit Acession No.	AB037816.1	4F112152.1	AA628056.1	İ	387675.1	V95708.1	AL163246.2	A95929.1	3F526041.1	4504374 NT	7662027 NT	7662027 NT	7507540	710/	3E393873.1	5453665 NT	5453665 NT	VF240786.1	4758423 NT	4504374 NT	(F015452.1	7705800 NT	T705900 NT	\B037807.1	Al290264.1	7657185 NT	7661979 NT	A447931.1	VA447931.1	7705764 NT	7705764 NT
Most Similar (Top) Hit BLAST E Value	8.0E-95	8.0E-95	8.0E-95	7.0E-95	7.0E-95	7.0E-95	7.0E-95	7.0E-95	3.0E-95	2.0E-95	2.0E-95	2.0E-95	30 10 0	2.0E-30	2.0E-95 E	2.0E-95	2.0E-95	2.0E-95	2.0E-95	2.0E-95	2.0E-95	2.0E-95	2.0E-95	2.0E-95	2.0E-95	2.0E-95	2.0E-95	2.0E-95	2.0E-95	2.0E-95	2.0E-95
Expression Signal	2.83	2.41	89'8	9.46	9.48	5.94	1.38	1.03	1.76	0.86	1.6	1.6	7 70	1.10	3.3	1,3	1.3	16.55	2.46	0.88	3.51	2.78	2.78	96.0	1.02	2.3	2.57	0.98	0.98	3.69	3.69
ORF SEQ ID NO;		28236		20058	20059			24623	25087	20694	21387	21388	24872	21012	21676	22147	22148	22191			22842	23232	23233	23268	23390	23940	24615			25121	25122
Exan SEQ ID NO:	17531	17987	19242	10240	10240	14169	14213	14857	15261	10847	11529	11529	44703	361	98/11	12236	12256	12294	12342	10848	13045	13433	13433	13479	13604	14163	14846	14895	14895	15287	15287
Probe SEQ ID NO:	7681	8088	6896	274	274	4270	4316	4982	5340	922	1625	1625	1807	3 8	ODAL	23/6	2376	2417	2468	2787	3120	3517	3517	3565	3690	4564	4971	6022	5022	5367	5367

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					TT1)														Ī	Ī	Ī			:	7								
Top Hit Descriptor	Human muscle-type phosphofructokinase (PFK-M) gene, exon 7	Homo sapiens ciliary dynein heavy chain 9 (DNAH9) mRNA, complete cds	Homo saplens huntingtin (Huntington disease) (HD), mRNA	Homo sapiens bone morphogenetic protein receptor, type IA (BMPR1A) mRNA	Homo sapiens glutathione S-transferase theta 2 (GSTT2) and glutathione S-transferase theta 1 (GSTT1)	genes, complete cds	Homo sapiens adenylosuccinate lyase (ADSL), mRNA	zt23h04.r1 Soares ovary tumor NbHOT Homo lapiens cDNA clone IMAGE:714007 5' similar to TR:G1087084 G1067084 F55H2.6;	223h04.r1 Soares ovary tumor NbHOT Homo capiens cDNA clone IMAGE:714007 5' similar to	DOS ENIOSAS 2000/004 FOUNDED I	RCG-FN0019-280500-011-G11 FN0019 Homo, sapiens cDNA	RC6-FN0019-290600-011-G11 FN0019 Homo sapiens cDNA	601437232F1 NIH_MGC_72 Homo sapiens cDNA clone IMAGE:3922423 5'	601497608F1 NIH_MGC_70 Homo sepiens cDNA clone IMAGE:3899761 5'	601497608F1 NIH_MGC_70 Homo saplens cDNA clone IMAGE:3899761 5'	PM0-LT0019-090300-002-409 LT0019 Homo sapiens cDNA	Homo sapiens chromosome 21 unknown mRNA	MR0-HT0559-250200-002-d07 HT0559 Homo sapiens cDNA	Homo saplens chromosome 21 segment HS21C001	Human glyceraldehyde-3-phosphate dehydrogenase pseudogene 3'end	Homo sapiens KIAA0763 gene product (KIAA0763), mRNA	Homo sapiens KIAA0763 gene product (KIAA0763), mRNA	Homo capiens myosin, heavy polypeptide 2, skeletal muscle, adult (MYH2), mRNA	Homo sapiens mRNA for KIAA1172 protein, pertial cds	Homo sapiens mRNA for KIAA1172 protein, partial cds	Homo sapiens mRNA for KIAA1172 protein, pertial cds	Homo sapiens phosphodiesterase 6A, cGMP-specific, rod, alpha (PDE6A), mRNA	Homo sapiens transient receptor potential channel 5 (TRPC5), mRNA	H.sapiens DNA for monoamine oxidase type A (7) (partial)	Homo sapiens A kinase (PRKA) anchor protein 1 (AKAP1), mRNA	Homo sapiens A kinase (PRKA) anchor protein 1 (AKAP1), mRNA	Human type IV collagenase (CLG4B) gene, excn 5	Human type IV callagenase (CLG4B) gene, exch 5
Top Hit Database Source	LN TA	NT	TN	FN	<u> </u>	Ę	NT	EST HUMAN	FOT LIMANI	NAMON TO FOL	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	EST_HUMAN	TN	NT	NT	NT	NT	LN T	NT	NT	LN	NT	NT	NT	NT	NT	۲
Top Hit Acession No.	A59724.1	4F257737.1		4757853 NT		-2407	11418164 NT	1.0E-95 AA284651.1			Ì				8.0E-98 BE907607.1						7662289 NT	7662289 NT	TN 653939 NT		5.0E-96 AB032998.1	\B032998.1	11416767 NT	6912735 NT	60812.1	11424399 NT	11424399 NT		168347.1
Most Similar (Top) Hit BLAST E Value	2.0E-95 M	2.0E-95[AI	2.0E-95	2.0E-95		2.0E-95 AF	2.0E-95	1.0E-95	4 00 05	1.05-93	1.05-95	1.0E-95	9.0E-96	8.0E-96	8.0E-98	8.0E-96	7.0E-96	6.0E-96	6.0E-98	6.0E-98 M26873.1	6.0E-96	6.0E-96	6.0E-98	5.0E-96	5.0E-96	5.0E-98	5.0E-98	5.0E-96	5.0E-96 X60812.1	5.0E-98	5.0E-96	5.0E-98 M68347.1	5.0E-96 M68347.1
Expression Signal	4.64	2.25	1.82	2.36		1.98	4.34	7.73	7 73	20,	4.85	4.85	1.67	0.82	0.82	2.66	0.95	0.85	96'0	26.15	1.98	1.98	2.09	2.7	3.08	3.06	2.31	0.98	1.22	4.23	4.23	1.81	1.81
ORF SEQ ID NO:	25748			28245		25285		25427	25,730	20150	26460	26461			20202	_	23538		22897	23159	28927	28928	28972	20098	20599	20600		22713			26194	26788	26789
Exon SEQ ID NO:				17996		١	19338	15371	15274	1000	16298	16298	16846	12666	12666				13197			18644	18682	10279	10751	10751	12447	12919	14694	16048	16048		16599
Probe SEQ ID NO:	5734	5957	6055	8106		9452	9840	5450	27.50	2020	5437	6437	6767	435	435	5383	3834	2213	3276	3437	8831	8831	8870	317	824	824	2570	2991	4810	6065	909	6719	6719

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Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
4097	13997		6.22	3.0E-96	H68656.1	EST_HUMAN	yr87h12.r1 Soares fetal liver spleen 1NFLS Homo saplens cDNA clone IMAGE:212327 5'
409	10355		3.49	2.0E-96	4503098 NT	NT	Homo sapiens chondroitin sulfate proteoglycan 4 (melanoma-associated) (CSPG4), mRNA
730	10662	20494	1.56	2.0E-96	AL163248.2	ΙN	Homo sapiens chromosome 21 segment HS21,0048
4651	14537	24326	1.58	2.0E-96	BE148074.1	EST_HUMAN	RC3-HT0230-040500-110-g02 HT0230 Homo sapiens cDNA
7191	17068		5.08	2.0E-98	AV689461.1	EST_HUMAN	AV689461 GKC Homo saplens cDNA clone GKCFMD07 5'
9151	18902		2.05	2.0E-96	AW249440.1	EST_HUMAN	2819351.5prime NIH_MGC_7 Homo sapiens cDNA clone IMAGE:2819351 5'
655	10590	20408	1.69	1.0E-96	Y18890.1	LN	Human endogenous retrovirus type K (HERV-K), gag, pol and env genes
1742	11643	21510	2.03	1.0E-98	AW955054.1	EST_HUMAN	EST367124 MAGE resequences, MAGC Homo saplens cDNA
1742	11643	21511	2.03	1.0E-98	AW955054.1	EST_HUMAN	EST367124 MAGE resequences, MAGC Homo sapiens cDNA
1806	11703	21580	0.89	1.0E-98	4503756 NT	LN	Homo sapiens flavin containing monooxygenass 2 (FMO2) mRNA
1806	11703	21581	0.89	1.0E-96	-	LN	Homo sapiens flavin containing monooxygenase 2 (FMO2) mRNA
2181	12068		1.33	1.0E-96	_	LΝ	Human hepatocyte growth factor gene, exon 1
2181			1.33	1.0E-96	M75967.1	TN	Human hepatocyte growth factor gene, exon 1
2219	12650	52009	1.88	1.0E-96	U51472.2	INT	Felis catus superfast myosin heavy chain (sMy∤HC) mRNA, complete cds
							Homo sapiens similar to ectonucleotide pyrophysphatase/phosphodlestarase 3 (H. sapiens) (LOC63214),
7058	16935	27125	20.65	1.0E-96	11419429 NT	ΙN	mRNA
7138	17015	27208	1.98	1.0E-98	AF274863.1	LN.	Homo sapiens secretory pathway component Sec31B-1 mRNA, alternatively spilced, complete cds
787			1.64	1.0⊑-98	AB033116.1	L'X	Homo sapiens mRNA for KIAA1290 protein, pertial cds
7843		27939	1.64	1.0E-98	AB033116.1	N	Homo sapiens mRNA for KIAA1290 protein, partial cds
3285	13206	23006	0.95	6.0E-97	BF245240.1	EST_HUMAN	601863712F1 NIH_MGC_57 Homo seplens cDNA clone IMAGE:4081202 5'
6429	16319		2.75	6.0E-97	BE141849.1	EST_HUMAN	IL5-HT0117-011099-004-D07 HT0117 Homo sapiens cDNA
6672	16552	26747	1.76	5.0E-97	AL043314.2	EST_HUMAN	DKFZp434N0323_r1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434N0323 5'
6735	16614	26804	10.79	5.0E-97	AA418026.1	EST HUMAN	zv97e12.s1 Soares_NhHMPu_S1 Homo sapiens cDNA clone IMAGE:767758 3' smilar to TR:G1304125 G1304125 PMS4 MRNA;
7578	<u></u>		2.76	5.0E-97	BF154912.1	EST_HUMAN	RC0-BT0812-250900-032-a09 BT0812 Homo saplens cDNA
8832	18645		1.87	5.0E-97	BE148597.1	EST_HUMAN	MR0-HT0241-150500-010-b02 HT0241 Homo sapiens cDNA
8832	18645	28930	18.1	20E-97	BE148597.1	EST_HUMAN	MR0-HT0241-150500-010-b02 HT0241 Homo saplens cDNA
924	10849	20697	1.28	4.0E-97	BE004436.1	EST_HUMAN	CM0-BN0106-170300-293-e06 BN0106 Homo sepiens cDNA
1868			1.08	4.0E-97		NT	Homo sapiens brefeldin A-inhibited guanine nucleotide-exchange protein 2 (BIG2), mRNA
6082	16027	26167	6.1	4.0E-97	Y11339.2	NT	Homo sapiens mRNA for GalNAc alpha-2, 6-slalytransferase I, long form
6082	16027		6.1	4.0E-97	Y11339	N	Homo sepiens mRNA for GalNAc alpha-2, 6-sialytransferase I, long form
6867		26939		4.0E-97		뉟	Homo sapiens v-src avian sarcoma (Schmidt-Ruppin A-2) viral oncogene homolog (SRC), mRNA
7328	17232		1.17	4.0E-97	AB011166.1	본	Homo sapiens mRNA for KIAA0594 protein, partial cds

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Table 4
Single Exon Probes Expressed in Heart

Top Hit Descriptor	Homo saplens mRNA for KIAA0594 protein, partial cds	Homo sapiens AXL receptor tyrosine kinase (AXL), transcript variant 1, mRNA	Homo sapiens AXL receptor tyrosine kinase (AXL), transcript variant 1, mRNA	Homo sapiens mRNA, similar to rat myomegalin, complete cds	Homo sapiens mRNA for KIAA1290 protein, pertial cds	Homo sapiens mRNA for KIAA1290 protein, pertial cds	Homo saplens G-2 and S-phase expressed 1 (GTSE1), mRNA	Homo sapiens mRNA for KIAA1172 protein, perital cds	Homo sapiens amyloid beta (A4) precursor protein (protease nexin-II, Alzheimer disease) (APP), mRNA	Homo sapiens amyloid beta (A4) precursor probein (protease nexin-II, Alzheimer disease) (APP), mRNA	Homo sapiens N-myc (and STAT) Interactor (NIMI), mRNA	Human beta-prime-adaptin (BAM22) gene, exon 7	Homo sapiens pericentrin (PCNT) mRNA	Homo sapiens eukaryotic translation elongation factor 1 alpha 1 (EEF1A1) mRNA	601339520F1 NIH_MGC_53 Homo sapiens cDNA done IMAGE:3681821 5'	Homo sapiens KIAA0649 gene product (KIAA0649), mRNA	Homo sapiens KIAA0649 gene product (KIAA0649), mRNA	nk29g02.s1 NCI_CGAP_Co11 Homo sapiens cDNA clone IMAGE:1014962 3'	Homo sapiens ribosomal protein S15 (RPS15); mRNA	Homo sapiens ribosomal protein S15 (RPS15); mRNA	PM4-BT0724-010400-008-a12 BT0724 Homo sapiens cDNA	Homo sapiens cat eye syndrome critical regionigene 1 (CECR1), mRNA	Homo sapiens death-associated protein (DAP), mRNA	Homo sapiens death-associated protein (DAP), mRNA	Human mRNA for amylold A4(751) protein	Homo sapians succinate CoA ligase, GDP-forming, alpha subunit (SUCLG1), mRNA	Homo sapiens mRNA for KIAA1365 protein, partial cds	Homo saplens mRNA for KIAA1005 protein, partial cds	Homo sapiens mRNA for KIAA1005 protein, partial cds	PM4-BT0724-010400-008-a12 BT0724 Homo sapiens oDNA	Homo sapiens hPAD-colony10 mRNA for peplidylarginine deiminase type I, complete cds	Homo sapiens IL2-Inducible T-cell kinase (ITK), mRNA
Top Hit Database Source	TN					NT		NT				TN			EST_HUMAN			EST_HUMAN			T_HUMAN				Z	NT ·	NT	NT	NT	T HUMAN	Ä	K
Top Hit Acession No.	4.0E-97 AB011166.1	11863122 NT	11863122 NT	4.0E-97 AB042557.1	AB033116.1	4.0E-97 AB033116.1	11418318 NT	3.0E-97 AB032998.1	4502166 NT	4502166 NT	4758813 NT	U36255.1	5174478 NT	4503470 NT	3E566488.1	11427757 NT	11427757 NT	4A553761.1	11426272 NT	11428272 NT	BE090973.1	8393092 NT	4758119 NT	4758119 NT	X06989.1	11321580 NT	9.0E-98 AB037786.1				AB033768.1	5031810 NT
Most Similar (Top) Hit BLAST E Value	4.0E-97	4.0E-97	4.0E-97	4.0E-97	4.0E-97	4.0E-97	4.0E-97	3.0E-97	3.0E-97	3.0E-97	3.0E-97	3.0E-97	3.0E-97	1.0E-97	1.0E-97 BE	1.0E-97	1.0E-97	1.0E-97 A	1.0E-97	1.0E-97	9.0E-98	9.0E-98	9.0E-98	9.0E-98	9.0E-98 XC	9.0E-98	86-30.6	9.0E-98	9.0E-98	9.0E-98	8.0E-98 A	8.0E-98
Expression Signal	1.17	1.76	1.76	15.68	2.31	2.31	3.83	1.17	10.96	10.96	1.77	1.92	1.14	12.55	2.19	3.41	3.41	282	13.54	13.54	8	1.29	4.74	4.74	2.77	241	1.39	2.24	2.24	4.97	0.89	1.04
ORF SEQ ID NO:	27434		28642	28124		28129		20028	20633	20634	21195		22948	24349		28229	28230	28768			20658	21012					27423		28498	20858	21112	21303
Exon SEQ ID NO:	17232	18376	18376	17882	17885	l	19012	10210	10783	10783	12694	L		14556		17979	17979	18494	17912	17912	10809	11162	16503	L	17148	17195	17223	18246		10809	11256	11444
Probe SEQ ID NO:	7328	8503	8503	8733	8736	8736	9331	243	856	928	1423	2389	3223	4670	5872	8088	8088	8629	8763	8763	883	1255	6623	6623	7271	7319	7355	8369	8369	9346	1350	1540

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Top Hit Descriptor	Homo sapiens IL2-Inducible T-cell kinase (ITK), mRNA	Homo sapiens PMS2L16 mRNA, partial cds	Homo saplens PMS2L16 mRNA, partial cds	Human mitochondrial creatine kinase (CKMT) gene, complete cds	Homo sapiens chromosome 21 segment HS21,0001	ht68f02xf NCI_CGAP_Lu24 Homo saplens cDNA clone IMAGE:3151899 3'	AJ403124 3.4 (downregulated in larynx carcinoma) Homo sapiens cDNA clone I8	Homo sapiens mRNA for KIAA0707 protein, pertial cds	7B18H01 Chromosome 7 Fetal Brain cDNA Library Homo saplens cDNA clone 7B18H01	Homo sapiens activator of S phase kinase (ASIK), mRNA	Homo sapiens activator of S phase kinase (ASK), mRNA	yo17g09.r1 Soares adult brain N2b5HB55Y Homo sapiens cDNA clone IMAGE:178240 5'	AJ403124 3.4 (downregulated in larynx carcinoma) Homo sapiens cDNA clone i8	AJ403124 3.4 (downregulated in larynx carcinoma) Homo sapiens cDNA clone I8	Human fumarasa precursor (FH) mRNA, nuclear gene encoding mitochondrial protein, complete cds	Homo sapiens Ran GTPase activating protein 1 (RANGAP1), mRNA	601172658F1 NIH_MGC_17 Homo sepiens cDNA clone IMAGE:3528134 5	Homo seplens chromosome 21 segment HS21,0002	Homo sapiens potassium channel subunit (HERG-3) mRNA, complete cds	Homo sapiens fatty-acid-Coenzyme A ligase, Icng-chain 4 (FACL4) mRNA	Homo saplens attractin precursor (ATRN) gene, exon 16	Homo sapiens attractin precursor (ATRN) genė, exon 16	Homo sapiens low density lipoprotein receptor related protein-deleted in tumor (LRPDIT), mRNA	Homo saptens low density lipoprotein receptor related protein-deleted in tumor (LRPDIT), mRNA	Homo sapiens protein tyrosine kinase 2 beta (FTK2B) mRNA	Homo sapiens PDZ domain-containing guanine nucleotide exchange factor i (LOC51735), mRNA	Homo sapiens SH3-domain GRB2-like 2 (SH3GL2), mRNA	Homo saplens SH3-domain GRB2-like 2 (SH3GL2), mRNA	H.saplens arginase gene exon 3 (EC 3.5.3.1)	Homo sapiens AIM-1 protein (LOC51151), mRNA	Homo sepiens chromosome 12 open reading frame 3 (C12ORF3), mRNA	tw36b04.x1 NCI_CGAP_Ut1 Homo sapiens cDNA olone IMAGE:2261743 3' similar to SW:RL2B_HUMAN P29316 60S RIBOSOMAL PROTEIN L23A.;
Top Hit Database Source	M	IN	TN	ΙN	TN	EST_HUMAN	EST_HUMAN	ᅜ	EST_HUMAN	NT	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	TN	Ę	EST_HUMAN	LΝ	LN	NT.	TN	LN	LN	N	LN	TN	TN	TN	NT	NT	LN	EST_HUMAN
Top Hit Acession No.	5031810 NT	4B017007.1	AB017007.1	J04469.1	8.0E-98 AL163201.2	BE348727.1	3.0E-98 AJ403124.1	3.0E-98 AB014607.1	AA077498.1	11419210 NT	. 11419210 NT	H46698.1	3.0E-98]AJ403124.1	3.0E-98 AJ403124.1	J59309.1	11418177 NT	BE294281.1	2.0E-98 AL163202.2	4F032897.1	4758331 NT	2.0E-98 AF218902.1	4F218902.1	8055269 NT	8055269 NT	4758975 NT	7706512 NT	11428813 NT	11428813 NT	X12664.1	7705868 NT	11435947 NT	41882007.1
Most Similar (Top) Hit BLAST E Value	80E-98	8.0E-98	8.0E-98	8.0E-98				3.0E-98	3.0E-98	3.0E-98	3.0E-98	3.0E-98		3.0E-98	3.0E-98	3.0E-98	2.0E-98 B	2.0E-98	2.0E-98	2.0E-98	2.0E-98	2.0E-98 A	2.0E-98	2.0E-98	2.0E-98	2.0E-98	2.0E-98	2.0E-98	2.0E-98 X12664.1	2.0E-98	2.0E-98	1.0E-98
Expression Signal	1.04	0.98	96.0	5.03	0.88	1.29	1.21	1.85	2.09	1.7	1.7	3.31	1.6	1.6	5.15	2.47	29.05	1.45	96.0	4.94	1.51	1.51	6:39	6:39	1.09	4.66	3.87	3.87	1.6	1.18	1.43	18.93
ORF SEQ ID NO:	21304	21468		23424				22329		26169	26160				28448		21816	21983					24711	24712	24795	25028	27073	27074	27672		25301	20172
Exan SEQ ID NO:	L	11597		13638	14921	19258		12436	12582	16020	16020	16960		17535	18199	18373	11924	12078	14099	L.		14608	14939	14939	15029	15224			17367	17801	19026	10345
Probe SEQ ID NO:	1640	1695	1695	3726	5049	9717	2131	2565	2720	6147	6147	7083	7685	7685	8322	9895	2033	2191	4189	4244	4720	4720	6909	5069	5163	2303	7004	7004	7497	7951	9350	399

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acesslon No.	Top Hit Database Source	Top Hit Descriptor
447	10391	20212	2.38	1.0E-98 A	AW998611.1	EST_HUMAN	PM0-BN0065-100300-001-c06 BN0065 Homo sapiens cDNA
1756	11655	21528	16.96		1.0E-98 N49818.1	EST_HUMAN	yv23f05.r1 Soares fetal liver spleen 1NFLS Hoino sapiens cDNA clone IMAGE:243585 5' similar to PIR:S54204 S54204 ribosomal protein L29 - human ;
							zp98c09.r1 Stratagene muscle 937209 Homo sapiens cDNA clone IMAGE:628240 5' similar to TR:G806562
5258	_	24953	6.96	١	1.0E-98 AA195854.1	EST_HUMAN	G806562 NEBULIN.;
7203		27268	1.36		1.0E-98 AF141349.1	TN	Homo sapiens beta-tubulin mRNA, complete cds
7203	17080	27267	1.38		AF141349.1	NT.	Homo sapiens beta-tubulin mRNA, complete cits
5681	15590	25692	4.29	9.0E-99	9.0E-99 AW968635.1	EST_HUMAN	EST380711 MAGE resequences, MAGJ Homo sapiens cDNA
8465	18338	28602	2.6		9.0E-99 AI479829.1	EST_HUMAN	tm69h07.x1 NCI_CGAP_BIn25 Homo sapiens cDNA clone IMAGE:2163421 3' similar to SW:BID_HUMAN P55957 BH3 INTERACTING DOMAIN DEATH AGONIST
8465	18338	28603	2.6		9.0E-99 AI479829.1	EST_HUMAN	tm69h07.x1 NCI_CGAP_Brn25 Homo saplens cDNA clone IMAGE:2163421 3' similar to SW:BID_HUMAN_P58957 BH3 INTERACTING DOMAIN DEATH AGONIST;
8716	18533	28817	1.84	9.0E-99	9.0E-99 AA134604.1	EST HUMAN	zл90d02.r1 Strategene lung carcinoma 937218 Homo sapiens cDNA clone IMAGE:565443 6' similar to TR:С662994 G682994 GPI-ANCHORED PROTEIN P137.;
2902	16942	27134	1.19	8.0E-99	9635487 NT	N T	Human endogenous retrovirus, complete genome
5561	18477	25550	9.2	7.0E-99	7.0E-99 AF035808.1	LZ LZ	Homo saplens oscillin (hLn) gene, exon 5
8884		28988	2.31	7.0E-99 AF	AF001886.1	Z	Homo sapiens NK-receptor (KIR-G2) gene, linker region exon
2080		21863	0.93	6.0E-99	11430555 NT	NT	Homo sapiens cysteine-rich repeat-containing protein S52 precursor, (LOC51232), mRNA
2080		21864	0.83	6.0E-99		LN.	Homo saplens cysteine-rich repeat-containing protein S52 precursor, (LOC51232), mRNA
3814	13728	23517	1.87	6.0E-99]A\	AW976364.1	EST_HUMAN	EST388473 MAGE resequences, MAGN Homo saplens cDNA
4842	14530	24318	1.16	6.0E-99	4502660 NT	L	Homo saplens CD34 antigen (CD34) mRNA
6013		26047	2.36	6.0E-99	6.0E-99 L43610.1	NT	Homo saplens polycystic kidney disease (PKDi) gene, exons 27-30
6013	i	26048	2.36	6.0E-99	6.0E-99 L43610.1	NT	Homo sapiens polycystic kidney disease (PKD _i i) gene, exons 27-30
6718		26787	1.21	6.0E-99	6.0E-99 X99101.1	NT	H.sapiens mRNA for estrogen receptor
7089		27160	2.18	6.0E-99	6.0E-99 AB036429.1	NT	Homo sapiens NDST4 mRNA for N-deacet/lase/N-sulfotransferase 4, complete cds
7143		27213	3.57	6.0E-99 AF	AF080255.1	NT	Homo sapiens lodestar protein mRNA, complete cds
7143		27214	3.57	6.0E-99 AI	-0802	NT	Homo sapiens iodestar protein mRNA, complete cds
8102		28241	3.72	6.0E-99	11526299 NT	NT	Homo saplens BH3 interacting domain death agonIst (BID), mRNA
805		20871	98.0	6-30.5	5.0E-99 U35464.1	NT	Human protein C inhibitor (PCI-B) mRNA, complete cds
905		20672	0.86	5.0E-99	5.0E-99 U35464.1	NT	Human protein C inhibitor (PCI-B) mRNA, complete cds
1922		21696	, 2.36	5.0E-99 Y1	Y11365.1	NT	H.sapiens IMPA gene, exon 8
4463		24148	1.35	5.0E-99 AI	AF009660.1	NT	Homo sapiens T cell receptor beta locus, TCRBV7S3A2 to TCRBV12S2 region
9909	\perp	24709	2.48	5.0E-99	4758697 NT	NT	Homo sapiens mannosidase, alpha, class 2A, member 1 (MAN2A1), mRNA
9360	19032		2.1	5.0E-99	5.0E-99 BE890177.1	EST_HUMAN	801513157F1 NIH_MGC_71 Homo saplens dDNA clone IMAGE:3914391 5'

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Top Hit Descriptor	Human E2A/HLA fusion protain (E2A/HLF) mRNA, complete cds	xp09e06x1 NCI_CGAP_HN9 Homo sapiens cDNA clone IMAGE:2739874 3' similer to gb:M31212 MYOSIN LIGHT CHAIN ALKALI, NON-MUSCI_E ISOFORM (HUMAN);	Human Ku (p70/p80) subunit mRNA, complete cds	Homo sapiens short chain L-3-hydroxyacyl-CoA dehydrogenase precursor (HADHSC) gene, nuclear gene encoding mitochondrial protein, complete cds	zb46d08.r1 Soares, fetal_lung_NbH119W Homo sepiens cDNA clone IMAGE:306635 5' similar to gb:M16182 BETA-GLUCURONIDASE PRECURSOR (HUMAN);	Homo sapiens myosin X (MYO10) mRNA, complete cds	Homo sapiens intersectin long isoform (ITSN) mRNA, complete cds	Homo sapiens GA-binding protein transcription factor, alpha subunit (60kD) (GABPA), mRNA	Human Ku (p70/p80) subunit mRNA, complete cds	Homo sapiens truncated Niemann-Pick C3 protein (NPC3) mRNA, complete cds	Homo sapiens truncated Niemann-Pick C3 protein (NPC3) mRNA, complete cds	Homo sapiens FK506-binding protein 6 (36kD) (FKBP6) mRNA, and translated products	Homo saplens FK506-binding protein 6 (36kD) (FKBP6) mRNA, and translated products	Human interferon-alpha receptor (HuIFN-alpha-Rec) mRNA, complete cds	Homo sapiens fatty acid amide hydrolase (FAẬH) gene, exon 14	Homo sapiens fatty acid amide hydrolase (FAAH) gene, exon 14	Homo sapiens chromosome 21 segment HS21C081	Homo sapiens ALEX1 protein (LOC51309), mRNA	hd02h02x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2908371 3' similar to TR:002711 002711 PRO-POL-DUTPASE POLYPROTEIN ;	Homo sapiens mRNA for KIAA1005 protein, partial cds	Homo sapiens glutathione S-transferase theta!2 (GSTT2) and glutathione S-transferase theta 1 (GSTT1)	ad Ico complete cas	Homo saplens chromosome 21 segment HS21C047	Homo sapiens chromosome 21 segment HS21C047	Homo saplens Testis-specific XK-related protein on Y (XKRY), mRNA	Homo sapiens Testis-specific XK-related protein on Y (XKRY), mRNA	xv78b11.x1 NCI_CGAP_Brn53 Homo saplens cDNA clone IMAGE::2824605 3'	Homo sapiens chromosome 21 segment HS210006	Homo capiens chromosome 21 segment HS21C049	EST02975 Fetal brain, Stratagene (cat#838206) Homo sapiens cDNA clone HFBCR32
Top Hit Database Source	E	EST_HUMAN L	Г	T B	EST HUMAN	Т			TN TN	TN T	TN TN			t E	TN TN	F	F		EST HUMAN	Г	<u> </u>			TN T			EST_HUMAN x	TN T		EST_HUMAN E
Top Hit Acession No.	M95586.1	AW274792.1	M30938.1	AF095703.1		7	AF114487.1	11528150 NT	M30938.1		AF192523.1	4503730 NT	4503730 NT	J03171.1	AF098018.1	AF098018.1	AL163281.2	11419721 NT	AW340174.1	AB023222.1	1 00207034	Ī	7	AL163247.2	11418230 NT	11418230 NT	AW275237.1		.2	T05087.1
Most Similar (Top) Hit BLAST E Value	3.0E-99 N	2.0E-99 A	2.0E-99 N	2.0E-99				1.0E-99	1.0E-99	1.0E-99	1.0E-99	1.0E-99	1.0E-99	1.0E-99		1.0E-99	1.0E-99	1.0E-99	1.0E-99	1.0E-99	700 20 7		_	1.0E-100	1.0E-100	1.0E-100	1.0E-100			1.0E-100
Expression Signal	5.37	3.46	1.08	3.15	9.75	3.83	1.53	1.04	2.11	1.47	1.47	1.1	1.1	0.89	2.74	2.74	1.18	1.15	1.68	2.4	94.0	9.70	1.13	1.93	1.48	1.48	1.52	1.16	1.01	2.43
ORF SEQ ID NO:			22947	24127	27121	28583	20093	20152	21168	21298	21299	21858	21657	22766		13961			27657	28854			19780	19780	19859	19860	19883			20114
Exon SEQ ID NO:	16723	11128	13144	14337		18324		10329	11308	11441	11441	11781	11781	12973		14182	15063	17181	17353	18571		1	6866	6866	10048	10048	10065			10299
Probe SEQ ID NO:	6844	1219	3220	4443	7053	8451	312	375	1401	1537	1537	1885	1885	3046	4283	4283	5200	7305	7483	8683	24.06	8150		2	62	62	81	162	314	340

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Table 4
Single Exon Probes Expressed in Heart

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Single Extri Flobes Expressed in near	Top Hit Descriptor	Homo sapiens X-linked anhidroitic ectodermal/dysplasia protein gene (EDA), exon 2 and flanking repeat regions	Gigorilla DNA for ZNF80 gene homolog	RC3-HT0625-040500-022-b09 HT0625 Homo sapiens cDNA	Homo saplens DKFZP588M0122 protein (DKFZP586M0122), mRNA	Homo saplens DKFZP586M0122 protein (DKF-ZP586M0122), mRNA	UI-H-BI1-afk-c-07-0-UI.s1 NCI_CGAP_Sub3 Homo sapiens cDNA clone IMAGE:2722164 3'	qf62f09.x1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1754633 3' similar to SW:CYT_COTJA P81061 CYSTATIN;	Rat mRNA for short type PB-cacherin, complete cds	H.sapiens mRNA for IFN-gamma (pKC-0)	Homo saplens KIAA0957 protein (KIAA0957), mRNA	Homo sapiens RGH2 gene, retrovirus-like element	Homo sapiens myotubularin-related protein 1a mRNA, partial cds	Homo sapiens follicle stimulating hormone receptor (FSHR) mRNA	Homo sapiens small optic lobes (Drosophila) homolog (SOLH) mRNA	Homo sapiens small optic lobes (Drosophila) homolog (SOLH) mRNA	601863164F1 NIH_MGC_57 Homo saplens cDNA clone IMAGE:4080999 5'	AU118182 HEMBA1 Hamo sapiens cDNA clone HEMBA1003048 5'	Homo saplens NF-E2-related factor 3 gene, complete cds	AU140214 PLACE2 Hamo sapiens cDNA clane PLACE2000137 5'	yf38c08.s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:129134 3'	MR1-TN0046-060900-004-b05 TN0046 Homo sepiens cDNA	MR1-TN0048-060900-004-b05 TN0048 Homb saplens cDNA	Human mRNA for kidney epidermal growth factor (EGF) precursor	601647357F1 NIH_MGC_61 Homo saplens cDNA clone IMAGE:3931310 5	Homo sapiens chromosome 21 segment HS21 C003	Homo saplens mRNA for KIAA1485 protein, partial cds	w/37g09.x1 NCI_CGAP_Pr28 Homo sapiens ¿DNA clone IMAGE:2489920 3' similar to contains element	MER22 repetitive element;	PM0-BN0065-100300-001-c08 BN0065 Homp sapiens cDNA	Homo sepiens mRNA for KIAA1626 protein, partial cds	Homo sapiens mRNA for KIAA1626 protein, partial cds	hh83c11.y1 NCI_CGAP_GU1 Homo sapiens cDNA clone IMAGE:2969396 5
פול באסוו בוסק	Top Hit Database Source	Þ	N	EST_HUMAN	TN	NT	EST_HUMAN	EST HUMAN	N	N	IN	IN	IN	IN	LN	LN	EST_HUMAN	EST_HUMAN	NT	EST_HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	LN	EST_HUMAN	ΙN	NT		EST_HUMAN	EST_HUMAN	LN	NT	EST_HUMAN
3 110	Top Hit Acession No.	\F003528.1	(89631.1	3E180609.1	7661685 NT	7661685 NT	W 207555.1	AI200857.1	383349.1	(62468.1	11418976 NT	J11078.1	AF057354.1	4503792 NT	5032104 NT	5032104 NT	3F244218.1	4U118182.1	4F135116.1	AU140214.1	310887.1	3F376478.1	3F376478.1	K04571.1	3F103853.1	4L163203.2	AB040918.1		41972388.1	4W998611.1	AB046846.1	AB046846.1	AW630487.1
	Most Similar (Top) Hit BLAST E Value	1.0E-100	\sim	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100		•	1.0E-100	-		1.0E-100
	Expression Signal	1.53	7.04	4.17	2.43	2.43	1.33	1.46	1.45	1.08	1.87	245	1.49	1.87	3.07	3.07	1.8	1.4	1.55	5.1	1.41	1.4	1.4	6.99	6.19	6.44	3.2		1.53	1.67	1.73	1.73	1.69
	ORF SEQ ID NO:			20255		20764		21294		22160	22429		23786	23816	24665	24666		25478	25496	25946	26049	26146	26147	26151	27033		27398			26625	27587		27740
	SEQ ID NO:	10375	10425	10442	10920	10920	11433	11437	12082	12267	12539	12913	14016	14043	14897	14897	15156	15415	15432	15821	15919	16010	16010	16013	16841	16860	17198			16439	17378	17378	17514
	Probe SEQ ID NO:	430	481	200	1002	1002	1528	1533	2195	2388	2674	2985	4116	4143	5024	5024	5232	5486	5514	5915	6015	6116	6116	6119	6963	6983	7322		7369	7426	7527	7527	7684

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	Exon ORF SEQ Expression (Top) Hit Acession Signal No: Signal Value Source	17614 27741 1.69 1.0E-100 AW 630487.1 EST_HUMAN INB3C11.71 NCI_CGAP_GU1 Homo sapiens cDNA clone IMAGE:2969396 5'	1.29 1.0E-100 Y10391.1 NT	28275 5.23 1.0E-100 BF327292.1 [EST_HUMAN	28745 2.14 1.0E-100 X94633.1 NT	NT	28831 4.56 1.0E-100 AF111170.3 NT	28832 4.56 1.0E-100 AF111170.3 INT	1.96 1.0E-100 AL163247.2 INT	186 1.0E-100 AF266285.1 NT Homo sapiens golgin-like protein (GLP) gene, complete cds	Homo sapiens glutathione S-transferase theta 2 (GS112) and glutathione S-transferase theta 2 (GS112) and glutathione S-transferase theta 1 (GS111) genes, complete cds	25302 2.21 1.0E-100 11646732 NT	25181 3.38 1.0E-100 11417974 NT	19871 1.22 1.0E-101 7110714 NT	19872 1.22 1.0E-101 7110714 NT	10605 20422 1.16 1.0E-101 AB007915.2 NT Homo sapiens mRNA for KIAA0446 protein, partial cds	10621 20447 4.45 1.0E-101 7110734 NT Homo sapiens ventral anterior homeobox 2 (VAX2), mRNA	7110734 NT	1.0E-101 1.32 1.0E-101 7657454 NT Home sepiens pescadillo (zebratish) homoiog 1, containing BKC1 domain (FES1), mKNA	Homo sapiens phosphoribosylglycinamide synthetase, phosphoribosylglyci	20679 3.44 1.0E-101 Z20856.1 NT	12.74 1.0E-101 BF681218.1 EST_HUMAN	10954 20796 1.63 1.0E-101 AI221878.1 EST_HUMAN qq99609.x1 Soares_NFL_T_GBC_S1 Homo(spplens cDNA clone IMAGE:1843335 3	1.0E-101 7662183 NT	1.0E-101 7662183 NT	21677 1.62 1.0E-101 4502996 NT	21792 1.6 1.0E-101 BE843070.1 EST_HUMAN	22080 1.66 1.0E-101 5729892 NT		2.56 1.0E-101 AJ237744.1 NT	1.0E-101 AJ237744.1 NT	12.14 1.0E-101 AJ252312.1 NT	13092 22897 2.51 1.0E-101 4885270 NT Homo sapiens germma-glutamytransterase 1 (1551 1) IIINNA
					١_		L			738			1						L						L								
-	Probe Exa SEQ ID SEQ NO: NC	7664 17	١.		L	1	L	1				1	┸	L	70 10	<u> </u>	688 10	<u> </u>	754 10			_	1036 10	1712 11	1712 11		┖	_	2572 12				3167 13

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Table 4
Single Exon Probes Expressed in Heart

	$\overline{}$	т-	_	7	_	_	$\overline{}$	_	т-	_	_	_	_		_	_		_	- -	Ŧ-	- -	i	1	7	—			-	τ	_			
Top Hit Descriptor	601458531F1 NIH_MGC_66 Homo sapiens cDNA done IMAGE:3862086 5	EST377629 MAGE resequences, MAGI Homo saplens cDNA	Homo sapiens RIBIIR gene (partial), exon 12	Homo sapiens RIBIIR gene (partial), exon 12	Homo sapiens ASH2L gene, complete cds, strillar to Drosophila ash2 gene	Homo sapiens butyrophilin, subfamily 2, member A1 (BTN2A1), mRNA	Homo sapiens butyrophilin, subfamily 2, member A1 (BTN2A1), mRNA	EST377212 MAGE resequences, MAGI Homo saplens cDNA	Homo sapiens cytoplasmic linker 2 (CYLN2), niRNA	Homo sapiens cytoplasmic linker 2 (CYLN2), niRNA	Homo sapiens Kruppel-type zinc finger protein (PEG3) mRNA, alternative splice form 4, partial cds	Homo sapiens Kruppel-type zinc finger protein (PEG3) mRNA, alternative splice form 4. partial cds	W65f12.x1 NCI_CGAP_Gas4 Homo sapiens cDNA clone IMAGE:2533487 3'	601109217F1 NIH_MGC_16 Homo sapiens d2NA clone IMAGE:3349901 5'	RC1-BT0313-220700-018-f12 BT0313 Homo sapiens cDNA	601764686F1 NIH_MGC_63 Homo sapiens cDNA clone IMAGE:3998837 5'	2K29g08.r1 Soares_pregnant_uterus_NbHPU Homo sapiens cDNA clone IMAGE:471998 5' similar to PIR:S54640 S54640 YD9336 03c nordain - verset	Human mRNA for pancreatic camma-dutamytransferase	Human mRNA for pancreatic gamma-glutamyltransferase	Homo saplens gamma-glutamyfiransferase 1 (GGT1), transcript variant 3, mRNA	601472808T1 NIH_MGC_68 Homo sapiens cDNA clane IMAGE:3876953 3'	601472808T1 NIH_MGC_68 Homo sapiens cDNA clone IMAGE:3875953 3'	Homo saplens Janus kinase 2 (a protein tyrosine kinase) (JAK2), mRNA	branched-chain alpha-keto acid dehydrogenase complex E1 alpha subunit [human, Genomic, 195 nt,	sagment 8 or 9.	Homo sapiens mRNA for KIAA0819 protein, partial cds	QV1-DT0068-240200-085-a01 DT0068 Homo sapiens cDNA	Homo sapiens chromosome 21 segment HS21C103	601108292F1 NIH_MGC_18 Homo saplens cDNA clone IMAGE:3344326 5'	Homo sapiens down-regulated in adenoma (DRA) mRNA	Human endogenous retroviral DNA (4-1), complete retroviral segment	Homo sapiens solute carrier family 2 (facilitated glucose transporter), member 9 (SLC2A9), mRNA	Homo sapiens solute carrier family 2 (facilitated glucose transporter), member 9 (SLC2A9), mRNA
Top Hit Database Source	EST_HUMAN	EST_HUMAN	ΙΝ	۲N	Ŋ	N	F	EST_HUMAN	Ņ	N	NT	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NAMI H TSH	LN	NT	NT	EST_HUMAN	EST_HUMAN	IN		Z	L	EST_HUMAN	⊥N	EST_HUMAN	NT	NT	NT	NT
Top Hit Acession No.	BF035327.1	AW965556.1	AJ237744.1	AJ237744.1	AB022785.1	5921460 NT	5921460 NT	AW965139.1	7427512 NT	7427512 NT	AF208970.1	AF208970.1	AW008475.1	BE257384.1	BF330759.1	BF029174.1	AA036800 1	X60069.1	X60069.1	9845492 NT	BE619667.1	BE619687.1	11429127 NT	, =0000	53832/.1	AB020626.1	AW939051.1	AL163303.2	BE252470.1	4557534 NT	M10976.1	11437148 NT	11437146 NT
Most Similar (Top) Hit BLAST E Value	1.0E-101			1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101		1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.05.101	-	-	1.0E-101	1.0E-101	1.0E-101	1.0E-101	_	-		1.0E-101	1.0E-102	-	1.0E-102	1.0E-102	1.0E-102	1.0E-102
Expression Signal	2.16	1.67	1.75	1.75	4.83	1.67	1.67	1.4	3.73	3.73	4.18	4.16	5.63	1.56	5.3	2.65	1.15	16,52	16.52	18.4	5.64	5.64	1.76	0	7.88	1.78	11.62	3.24	0.89	1.46	1.95	1.39	1.39
ORF SEQ ID NO:			22468	22469	23499	24609	24610	24954	25658	52669		26357	26441		26581	26764	27274	26850		27468	27694	27695	28040	70000	19897	28589		20111	20350	20526	20869	21002	21003
Exan SEQ ID NO:	Ш		12576	12678	13712		14840	15179	15583	15563	16196	16196		16330	16402	16573	17085	16458	16458	17263	17474	17474	17800	0,00	75101	18327	19189	10297	10540	10688	11017	11154	11154
Probe SEQ ID NO:	3203	3334	3354	3354	3800	4985	4965	5257	5851	5851	6333	6333	6416	6471	6544	6693	7208	7448	7446	7454	7623	7623	7950	C	7070	2 2	9610	88	8	758	1101	1247	1247

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Top Hit Descriptor	601299982F1 NIH_MGC_21 Homo sapiens cDNA done IMAGE:3628901 5'	em60c10.x1 Johnston frontal cortex Homo sapiens cDNA clone IMAGE:1539954 3' similar to SW:GG95_HUMAN Q08379 GOLGIN-95.;	ат60c10.x1 Johnston frontal cortex Homo sapiens cDNA clone IMAGE:1639954 3' similar to SW:GG95_HUMAN Q08379 GOLGIN-95.;	Homo saplens KIAA0187 gene product (KIAA0187), mRNA	AU141005 PLACE4 Homo sapiens cDNA clone PLACE4000650 5'	AU141005 PLACE4 Homo sapiens cDNA clone PLACE4000650 5	Homo sapiens chromosome 21 segment HS2(1007	601107843F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3343882 5'	y32c04.r1 Soares placenta Nb2HP Homo sapiens cDNA clone IMAGE:140934 5'	Homo sapiens protein phosphatase-1 regulatory subunit 7 (PPP1R7) gene, exon 7	Homo saplens HSC54 mRNA for heat shock cognate protein 54, complete cds	Homo sapiens histone deacetylase 7 (HDAC7), mRNA	Homo sapians histone deacetylase 7 (HDAC7), mRNA	ar82f09x1 Barstead colon HPLRB7 Homo sapiens cDNA clone IMAGE:2151785 3' similar to TR:Q13137	21915/ NUPOL. ;	Homo sapiens mRNA for Centaurin-alpha2 protein	AV710738 Cu Homo sapiens cDNA clone CuAAKD03 5'	QV3-NT0025-210600-236-h08 NT0025 Homo sapiens cDNA	601501107F1 NIH_MGC_70 Homo saplens cDNA clone IMAGE:3903145 5	AV694817 GKC Homo sapiens cDNA clone GKCEEE11 5'	AV694817 GKC Homo sapiens cDNA clone GKCEEE11 5'	Homo sapiens mRNA for KIAA0454 protein, partial cds	yd13d07.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:67021 5'	yd13d07,r1 Soares fetal liver spleen 1NFLS Flomo sapiens cDNA clone IMAGE:67021 5'	AU124629 NT2RM4 Homo seplens cDNA clone NT2RM4000309 5'	Homo sapiens myomesin (M-protein) 2 (165kJ) (MYOM2), mRNA	Homo sapiens myomesin (M-protein) 2 (165kD) (MYOM2), mRNA	RC-BT074-260499-014 BT074 Hamo saplens cDNA	RC-BT074-260499-014 BT074 Homo saplens cDNA	on57h04.s1 Soares_NFL_T_GBC_S1 Homo/sapiens cDNA clone IMAGE:1560823 3' similar to SW:CAV2_HUMAN P51638 CAVEOLIN-2. [1] ;	601439392F1 NIH_MGC_72 Homo saplens cDNA clone IMAGE:3924166 5'
Top Hit Database Source	EST_HUMAN	EST_HUMAN	EST HUMAN	N.	EST_HUMAN		F2	EST_HUMAN	EST_HUMAN	Ę	N	Ę	LN FA	100	EST HOMAN	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	Z	Z-	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN
Top Hit Acession No.	BE408447.1	1.0E-102 AI124669.1	N124669.1	7661979 NT	NU141005.1	\U141005.1	AL.163207.2	1.0E-102 BE251310.1	366488.1	₹5067133.1	\B034951.1	TN 865398 NT	7705398 NT		1.0E-102 AI459825.1		10710738.1	1.0E-102 BE763051.1	3E910555.1	1.0E-102 AV694817.1	V694817.1	AB007923.1	70383.1	70393.1	\U124629.1	11425430 NT	11425430 NT	N905037.1	N905037.1	14970786.1	1.0E-102 BE897468.1
Most Similar (Top) Hit BLAST E Value	1.0E-102	1.0E-102	1.0E-102	_	1.0E-102	1.0E-102	1.0E-102	1.0E-102	1.0E-102	1.0E-102	1.0E-102	1.0E-102	1.0E-102	L	1.0E-102/	1.0E-102/	1.0E-102	1.0E-102	1.0E-102	1.0E-102	1.0E-102		1.0E-102	1.0E-102	1.0E-102	1.0E-102	1.0E-102	1.0E-102 A	1.0E-102 A	1.0E-102	1.0E-102
Expression Signal	90.24	1.34	1.34	1.51	5.61	5.61	1.48	2.11	1.09	1.88	7.27	3.46	3.46	1	γ. γ.	8.58	2.53	4.2	2.53	1.36	1.36	4.08	1.52	1.62	3.58	2.03	2.03	2.9	2.9	2.3	2.38
ORF SEQ ID NO:	21162	22044	22045		22815	22816	23814	23994	24705	25025		25522	25523		١		26524	26857	26904	27014	27015	27069	27448	27449	27490	28052	28053	28070	28071		
Exan SEQ ID NO:	11303	12145	12145	12954	13021	13021	14039	14211	14933	15221	15433	15453	15453	J	ł		18354	16666	16711	16823	16823	16878	17243	17243	17282	17811	17811	17831	17831	l.	Н
Probe SEQ ID NO:	1398	2261	2261	3026	3094	3094	4139	4314	5063	5300	5515	5536	5536		2800	6351	6495	6787	6832	6945	6945	7001	7374	7374	7415	7981	7861	7981	7981	8002	8421

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Top Hit Descriptor	Homo sapiens UDP glycosyltransferase 2 family, polypeptide B11 (UGT2B11) mRNA	Homo sapiens UDP glycosyltransferase 2 family, polypeptide B11 (UGT2B11) mRNA	RC6-ET0072-150600-011-F01 ET0072 Homo sapiens cDNA	Human chromosome 16 creatine transporter (SLC6A8) and (CDM) paralogous genes, complete cds	Homo saplens chromosome 21 segment HS21C080	xk07c12.x1 NCI_CGAP_Co20 Homo sapiens cDNA clone IMAGE:2666038 3'	601500405F1 NIH_MGC_70 Homo sapiens cDNA done IMAGE:3802305 5	601500405F1 NIH_MGC_70 Homo saplens ¢DNA done IMAGE:3902305 5'	Homo sapiens mRNA for KIAA0235 protein, partial cds	Homo sapiens nucleolar protein (KKE/D repeat) (NOP56) mRNA	Homo sapiens mRNA for pregnancy-associated plasma protein-E (PAPPE gene)	601485388F1 NIH_MGC_69 Homo sapiens cDNA clone IMAGE:3887876 5	Homo sapiens phosphatidylinositol 4-kinase 230 (pi4K230) mRNA, complete cds	Homo sapiens smg GDS-ASSOCIATED PROTEIN (SMAP), mRNA	Homo saplens bone morphogenetic protein 8 (osteogenic protein 2) (BMP8) mRNA	Homo sapiens bone morphogenetic protein 8 (osteogenic protein 2) (BMP8) mRNA	AU134991 PLACE1 Homo sapiens cDNA clone PLACE1000965 5'	Homo sapiens promyelocytic leukemia zlnc finger protein (PLZF) gene, complete cds	yw91d08.s1 Soares_placenta_8to9weeks_2NbHP8to9W Homo sapiens cDNA clone IMAGE:259599 3'	601573113F1 NIH_MGC_9 Homo sepiens cDNA clone IMAGE:3834315 5'	UI-H-BW0-ajl-h-11-0-UI.s1 NCI_CGAP_Sub3 Homo sapiens cDNA clone IMAGE:2733165 3'	Homo saplens mRNA for KIAA1459 protein, partial cds	Macaca mulatta cyclophilin A mRNA, compleje cds	ab10d12.s1 Stratagene lung (#637210) Homo sapiens cDNA clone IMAGE:840407 3' similar to contains	The contract of the contract o	noine sapiens regional i (NKT I), microx	seq340 b4HB3MA-Cct109+10-Bio Homo sapiens cDNA clone b4HB3MA-Cct109+10-Bio-7 3	Homo sapiens septin 2 (SEP2) mRNA, partial cds	Homo sapiens glycine receptor alpha 2 subunit (GLRA2) gene, exon 4	tm58b05.x1 NCI_CGAP_Brn25 Homo sapieris cDNA clone IMAGE:2162289 3' similar to TR:Q13769 Q13769 ANONYMOUS.;
Top Hit Database Source	ΓN	TN	EST_HUMAN	۲	۲N	EST_HUMAN	EST_HUMAN	EST_HUMAN	N F	L	칟	EST_HUMAN	FN	N	N	Z	EST_HUMAN	TN	EST HUMAN	EST_HUMAN	EST_HUMAN	۲N	닐	MALTIN FOR	NAMOR STATE	- N	EST_HUMAN	NT	NT	EST_HUMAN
Top Hit Acession No.	4507822 NT	. 4507822 NT	BF359243.1	U41302.1	AL163280.2	AW300862.1	BE908158.1	BE908158.1	D87078.2	5453793 NT	AJ278348.1	BE877541.1	AF012872.1	7657592 NT	4502428 NT	4502428 NT	AU134991.1	AF060568.1	N32770.1	BE744722.1	AW 298245.1	AB040892.1	AF023861.1	A A 40 E E E 2 4	44.40000	114300/00/11	123683.1	AF178995.1	AF053490.1	AI590071.1
Most Similar (Top) Hit BLAST E Value	1.0E-102		1.0E-102	1.0E-102	1.0E-102	1.0E-102	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	_	1.0E-103	1.0E-103	1.0E-103	4 05 409		-	_		1.0E-103	1.0E-103
Expression Signal	1.89	1.89	2.78	4.04	2.82	4.77	0.82	0.82	8.1	0.83	1.01	7.29	2.32	1.04	1.27	1.27	1.57	2.22	0.88	2.43	3.44	1.08	2.41	1 18	7.10	80	3.02	1.72	5.37	1.68
ORF SEQ ID NO:	Ш	28554	28807	29051		25247	19861	19862	19894		20734	20985	21342	21643	21708	21709	22040	22175	22340		23058			20766		10407			26027	26069
Exon SEQ ID NO:	Ш		18525	18755	18837	19183	10049	10049	10078	10173	10888	11131	11482	11768	11827	11827	12141	12278	12449	12958	13253	13310	13609	13837	13057	0/001	_	15534	15903	16936
Probe SEQ ID NO:	8424	8424	8708	8947	9054	9601	ස	හ	63	201	865	1223	1578	1872	1932	1932	2257	2401	2578	3030	3333	3333	3692	3775	0757	10/0	3922	5919	5998	6033

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				_					_	_		-	******									
Top Hit Descriptor	tm88b05.x1 NCI_CGAP_Brn25 Homo sapien's cDNA clone IMAGE:2162289 3' similar to TR:Q13769 Q13769 ANONYMOUS.;	Homo sepiens dystrophin (muscular dystrophy, Duchenne and Becker types), includes DXS142, DXS164, DXS206, DXS230, DXS230, DXS230, DXS230, DXS269, DXS270, DXS272 (DMD), transcript varient Dp427m, mRNA	Homo saplens dystrophin (musculer dystrophy, Duchenne and Becker types), includes DXS142, DXS184, DXS206, DXS230, DXS230, DXS239, DXS269, DXS270, DXS272 (DMD), transcript varient Dp427m, mRNA	EST377849 MAGE resequences, MAGI Homo sapiens cDNA	601571537F1 NIH_MGC_55 Homo sapiens cDNA done IMAGE:3838545 5'	tm58b05.x1 NCI_CGAP_Bm25 Homo sapiens cDNA clone iMAGE:2162289 3' similar to TR:Q13769 Q13769 ANONYMOUS.;	tm58b05.x1 NCI_CGAP_Bm25 Homo sapien's cDNA clone IMAGE:2162289 3' similar to TR:Q13769 Q13789 ANONYMOUS.	EST27193 Human Brain Homo sapiens CDNA 5 end similar to None	AU140344 PLACE2 Homo sapiens cDNA clone PLACE2000374 5'	AU140344 PLACE2 Homo sapiens cDNA clone PLACE2000374 5'	7160e03.x1 Soares_NSF_F8_9W_OT_PA_P!_S1 Homo saptens cDNA clone IMAGE:3525964 3' similar to SW:PTNF_HUMAN Q16825 PROTEIN-TYROSINE PHOSPHATASE D1;	Homo sapiens triple functional domain (PTPRF interacting) (TRIO), mRNA	Homo sapiens triple functional domain (PTPRF interacting) (TRIO), mRNA	H.saplens mRNA for latent transforming growth factor-beta binding protein (LTBP-2)	EST375749 MAGE, resequences, MAGH Homo sapiens cDNA	au51g04.y1 Schneider fetal brain 00004 Homo sapiens cDNA clone IMAGE:2518329 5' similar to TR:O15045 O15046 KIAA0338 ;	olozdos ys NCI_CGAP_Lus Home sapiens cDNA clone IMAGE:1522283 5' similar to TR:Q62084 Q62084 PHOSPHOLIPASE C NEIGHBORING;	Homo sapiens NOD1 protein (NOD1) gene, exons 1, 2, and 3	Homo saplens NOD1 protein (NOD1) gene, exons 1, 2, and 3	AU136283 PLACE1 Homo sapiens cDNA clone PLACE1003923 5'	Homo saplens polycystic kidney disease (PKI)1) gene, exons 27-30	7e68a10.x1 Soares_NSF_F8_9W_OT_PA_P_S1 Homo saplens cDNA clone IMAGE:3287610 3' similar to contains MER29.t3 MER29 repetitive element;
Top Hit Database Source	EST HUMAN	Z	, Z	EST HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	NT	IN	EST_HUMAN	EST_HUMAN	EST HUMAN	N L	LN	EST_HUMAN	NT	EST_HUMAN
Top Hit Acession No.	AI590071.1	5032282 NT	5032282 NT	AW965776.1	BE748158.1	A1590071.1	AI590071.1	T31080.1	AU140344.1	AU140344.1	BF109244.1	6005921 NT	6005921 NT	Z37976.1	AW983676.1	AI878958.1	AI792759.1	AF149773.1	AF149773.1	AU136283.1	L43610.1	1.0E-103 BE644611.1
Most Similar (Top) Hit BLAST E Value	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103
Expression Signal	1.68	89:1	1.68	1.64	3.21	3.28	3.28	2.95	1.17	1.17	1.43	3.08	3.08	2.02	2.09	9.93	3.08	2.74	2.74	2.56	6.49	3.42
ORF SEQ ID NO:	26070			L	26361		26608			L	27118	27337	27338	27870	27905	27972	28250	28353	L		28136	29070
Exan SEQ ID NO:	15936	15102	15102	16153	16201	16426	16426	L	ı	16887	16927	17144	17144	17637	17665	17728	18004	18101	l _	18569	17892	18778
Probe SEQ IO NO:	6033	6092	6092	6289	6338	6568	6568	6823	7010	7010	7050	7287	7267	787	7815	7878	8115	8218	8218	8681	8743	8973

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Top Hit Database Source	Homo sapiens mannosidase, beta A, Iysosomal (MANBA) gene, and ubiquitin-conjugating enzyme E2D 3 NT (UBE2D3) genes, complete cds	NT Homo sepiens hypothetical protein FLJ20454 (FiL)20454), mRNA	NT Homo sapiens gene for AF-6, complete cds	EST_HUMAN DKFZp564H1072_r1 564 (synonym: hfbr2) Hoino sapiens cDNA clone DKFZp564H1072 5'	EST_HUMAN DKFZp564H1072_r1 564 (synanym: hlbr2) Hojno sepiens cDNA clone DKFZp564H1072 5'		zo22c06.s1 Stratagene cdon (#937204) Homo sapiens cDNA clone IMAGE:587626 3' similar to gb:Z14116_ma1 CD59 GLYCOPROTEIN PRE:CURSOR (HUMAN);	EST_HUMAN 601577460F1 NIH_MGC_9 Homo sepiens cDIVA clone IMAGE:3926438 5'	EST_HUMAN RC1-CT0249-110900-214-f12 CT0249 Homo sapiens cDNA	EST_HUMAN RC1-CT0249-110900-214-f12 CT0249 Homo sapiens cDNA		NT Human lymphocytic antigen CD59/MEM43 mRNA, complete cds	NT H.sapiens gene encoding phenylpyruvate tautomerase II	T_HUMAN	NT Homo sapiens mRNA for KIAA1276 protein, partial cds	NT Homo sapiens mRNA for KIAA1276 protein, partial cds	NT Homo sapiens mRNA for KIAA1172 protein, partial cds	NT Human mRNA for fibronectin (FN precursor)	NT Homo sapiens chromosome 21 unknown mRNA	NT Homo sepiens chromosome 21 unknown mRNA	NT Human Down Syndrome region of chromosomie 21 DNA	NT Human Dawn Syndrome region of chramosome 21 DNA	wjo3b12.x1 NCI_CGAP_Kid12 Homo sapiene cDNA clone IMAGE:2401727 3' similar to TR:Q14145 Q14145	wjo3b12.x1 NCI_CGAP_Kid12 Homo sapiens cDNA clone IMAGE:2401727 3' similar to TR:Q14145 Q14145 FST_HIMAN KIAA0132 PROTEIN, contains element LTR7 repetitive element;	HIMAN	HIMAN	LZ	EST HUMAN		
Top Hit Acession No.	1.0E-103 AF224669.1	11526291 NT	1.0E-103 AB011399.1		037549.3	4502428 NT			1.0E-104 BF334221.1	1.0E-104 BF334221.1	5031570 NT		1151.1	319436.1	1.0E-104 AB033102.1	1.0E-104 AB033102.1	-		1.0E-104 AF231920.1	1.0E-104 AF231920.1			_	68707 1	314182 1		25572	1 0E-104 BF448230.1	Γ	
Most Similar (Top) Hit BLAST E Value	1.0E-103	1.0E-103	1.0E-103	1.0E-104 ALI	1.0E-104 AL	1.0E-104	1.0E-104 /	1.0E-104	1.0E-104	1.0E-104	1.0E-104	1.0E-104 M34671.1	1.0E-104 Y1	1.0E-104	1.0E-104	1.0E-104	1.0E-104	1.0E-104 X02761.1	1.0E-104	1.0E-104	1.0E-104	1.0E-104 U43379.1	1.0E-104 AI7	1 OF-104 A17	1 OF-104	1 OF-104	105 104	1 0F-104	1.0E-104	1.0E-104
Expression Signal	1.72	2.65	2.21	2.6	2.8	1.81	7.16	1.91	1.15	1.16	1.55	7.64	2.82	1.54	68.0	66.0	0.91	4.28	6.0	6.0	1.33	1.33	8.46	8 48	1 50	1 52	3000	2.24	4,68	4.66
ORF SEQ ID NO:			25323			21617									23260	23261	23564	23957	24182	24183	25615		25919				l			
Exan SEQ ID NO:	18833	18855	18979	10202	1	11741	12035	L	L	1	12268			13265	13465	13465	13772	14179	14397	14397		L				Ł	L			1_
Probe SEQ ID NO:	9049	9079	9275	233	233	1845	2147	2157	2318	2318	2387	2842	2886	3345	3550	3550	3861	4280	4504	4204	5617	5617	5894	F8004	6073	200	2000	7288	7370	7370

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	Top Hit Descriptor	IL3-HT0819-080900-249-F07 HT0619 Homo sapiens cDNA	IL3-HT0819-080900-249-F07 HT0619 Homo sapiens cDNA	601581503F1 NIH_MGC_7 Homo sapiens cDNA done IMAGE:3935977 5'	601681603F1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:3935977 5'	AV728070 HTC Homo saplens cDNA clone H TCBYA07 5'	AU130765 NT2RP3 Homo sapiens cDNA clone NT2RP3001398 5'	Human beta4-integrin (ITGB4) gene, exons 19/20,21,22,23,24 and 25	RC0-HT0885-310700-021-b09 HT0885 Homo saplens cDNA	RC0-HT0885-310700-021-b09 HT0885 Homo sapiens cDNA	602141215F1 NIH_MGC_46 Homo sapiens cloNA clone IMAGE:4302507 5'	601312181F1 NIH_MGC_44 Homo sapiens cDNA clane IMAGE:3658676 5'	Homo sapiens amyloid beta (A4) precursor protein (protease nexin-II, Alzheimer disease) (APP), mRNA	Homo sapiens Mels1 (mouse) homolog (MEISI1) mRNA	Homo sapiens potassium channel subunit (HERG-3) mRNA, complete cds	Homo sapiens potassium channel subunit (HERG-3) mRNA, complete cds	Homo sapiens chromosome 21 segment HS2; C080	Human mRNA for KIAA0128 gene, partial cds	EST20609 Spleen I Homo sapiens cDNA 5' end similar to autoimmune antigen Ku, p70/p80 subunit	no10d05.s1 NCI_CGAP_Phe1 Homo sapiens cDNA clone IMAGE:1100265 3'	Homo sapiens 959 kb contig between AML1 and CBR1 on chromosome 21q22; segment 1/3	Homo sapiens bromodomain adjacent to zinc finger domain, 28 (BAZ2B), mRNA	Homo sapiens bromodomain adjacent to zinc finger domain, 2B (BAZ2B), mRNA	Homo sapiens dermatopontin (DPT), mRNA	EST373761 MAGE resequences, MAGG Homo sapiens cDNA	601445823F1 NIH_MGC_66 Homo saplens cDNA clone IMAGE:3850156 5	601445823F1 NIH_MGC_65 Homo sapiens cDNA clone IMAGE:3850156 5'	Homo saplens chromosome 21 segment HS21C008	Homo sapiens mRNA for KIAA0796 protein, partial cds	Homo saplens mRNA for KIAA0866 protein, complete cds	Homo saplens mRNA for KIAA0796 protein, partial cds	Homo saplens GTPase activating protein-like (GAPL), mRNA	Homo saplens GTPase activating protein-like (GAPL), mRNA	EST02975 Fetal brain, Stratagene (cat#936206) Homo sapiens cDNA clone HFBCR32
201 11000	Top Hit Database Source	Г	EST_HUMAN IL	EST_HUMAN 6	EST_HUMAN 6	EST_HUMAN A	EST_HUMAN A	노	EST_HUMAN R		EST_HUMAN 60	EST_HUMAN 60			F F	NT H	F	H	EST_HUMAN E	EST_HUMAN no					EST_HUMAN E	EST_HUMAN 6		T.	NT TN	H H	NT H			EST_HUMAN E
.G	Top Hit Acession No.	BF352841.1	BF352841.1	BE791713.1	BE791713.1	AV728070.1	AU130765.1	U66535.1	BE720191.1	BE720191.1	BF684288.1	BE393892.1	4502168 NT	4505150 NT	AF032897.1	AF032897.1	AL163280.2	D50918.1	AA318369.1	AA584808.1	AJ229041.1	7304922 NT	7304922 NT	11425532 NT	AW961688.1	BE868881.1	BE868881.1	AL163208.2	AB018339.1	AB020673.1	AB018339.1	11419196 NT	1419196	T05087.1
	Most Similar (Top) Hit BLAST E Value	1.0E-104	1.0E-104	1.0E-104		1.0E-104	1.0E-104	1.0E-104	1.0E-104	1.0E-104	1.0E-104	1.0E-104	1.0E-105	1.0E-105	1.0E-105	1.0E-105	1.0E-105	1.0E-105	1.0E-105	1.0E-105	1.0E-105	1.0E-105	1.0E-105	1.0E-105	1.0E-105	1.0E-105	1.0E-105	1.0E-105	1.0E-105	1.0E-105	1.0E-105	1.0E-105		1.0E-105
	Expression Signal	3.84	3.84	3.14	3.14	1.42	4.51	4.24	1.84	1.84	4.49	1.37	2.86	16.85	3.22	3.22	1.18	1.75	1.64	0.87	2.57	0.93	0.93	1.28	2.15	0.84	0.84	3.74	0.95	2.23	1.3	3.08	3.06	6.43
	ORF SEQ ID NO:	26630	26631	86872	27899	28037	28057	28110	28755	28756	28783		20082	19777	20323	20324	21553	21655	21928				23033		23689	24313	24314		24690	24740	24690	24840	24841	26754
	Exan SEQ ID NO:	16443	16443	17659		17797	17816	17864	18484	18484	18505	19340	12637	9866	10517	10517	11676	11780	12030	12554	12902	13228	13228	13508	13914	14528	14528	14735	14916	14965	14916	15124	15124	16559
	Probe SEQ ID NO:	7430	7430	2809	7809	7947	7966	8014	8617	8617	8841	9842	277	419	579	629	1777	1884	2142	2689	2875	3307	3307	3592	4008	4638	4638	4855	5044	2603	5161	6167	6167	6679

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			_	_	_		_	_	_	~	~	_			,		_2	inter.			**				*****	<u>-</u>	-		*	4
Top Hit Descriptor	ws50c10.x1 NCI_CGAP_Brn25 Homo sapiens cDNA clone IMACE:2500628 3' similar to SW:ACSA_PENCH P36333 ACETYL-COENZYME A SYNTHETASE	UI-H-Blop-abl-b-12-0-UI.s1 NCI_CGAP_Sub2 Homo sapiens cDNA clone IMAGE:27117823'	Homo sapiens SMARCA4 isoform (SMARCA4) gene, complete cds, alternatively spliced	Homo sapiens COL4A6 gene for a6(IV) collagen, exon 31	Homo sapiens Ran binding protein 11 (LOC51194), mRNA	wv74f07.x1 Soares_thymus_NHFTh Homo squiens cDNA clone IMAGE:2636301 3' similer to TR.:P87892 P87892 PROTEASE :	UI-HF-BN0-akt-g-07-0-UI.r1 NIH MGC 50 Homo sapiens cDNA done IMAGE:3078348 5'	tq79c01.x1 NCI_CGAP_Ut1 Homo sapiens cDNA clone IMAGE:2215008 3'	EST377629 MAGE resequences, MAGI Homo sapiens cDNA	Human dihydrofolate reductase pseudogene (psi-hd1)	Human dihydrofolate reductase pseudogene (psi-hd1)	Homo sapiens soluble neuropilin-1 mRNA, complete cds	Human epidermal growth factor receptor (EGFR) precursor-mRNA, exon 4, partial cds	ng41c05.s1 NCI_CGAP_Co3 Homo sapiens cDNA clone IMAGE:937352 3' similar to contains element LTR3 repatitive element;	ng41c05.s1 NCI_CGAP_Co3 Homo sapiens cDNA clone IMAGE:937352 3' similar to contains element	LIRS repetitive element;	Will safety of the concentration of the control of	Homo sapiens glutathione S-transferase theta[1 (GSTT1), mRNA	Homo saplens X-linked anhidrolitc ectodermal dysplasia protein gene (EDA), exon 2 and flanking repeat regions	Homo capiens sperm membrane protein BS-63 mRNA, complete cds	601149783F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:3502481 5'	q176h10x1 Soares_NhHMPu_S1 Homo saplens cDNA clone IMAGE:1878307 3'	Homo sapiens glutathione S-transferase theta 1 (GSTT1), mRNA	Homo sapiens glutathione S-transferase theta 1 (GSTT1), mRNA	Homo saplens mRNA for KIAA1326 protein, partial cds	Homo sapiens mRNA for KIAA1326 protein, partial cds	Homo sapiens hypothetical protein FLJ11273 (FLJ11273), mRNA	Homo saplens hypothetical protein FLJ11273 (FLJ11273), mRNA	Hamo saplens gene for activin receptor type IIB, complete cds	Homo sapiens mRNA for KIAA1278 protein, partial cds
Top Hit Database Source	EST HUMAN	EST_HUMAN	닏	N _T		EST HUMAN	Τ	Т	Г		IN	 LN	TN	EST HUMAN		Т	NIZWO L		<u> </u>		EST_HUMAN	EST_HUMAN C			±N	LN				± V
Top Hit Acession No.	AW007194.1	AW016879.1	AF254822.1	D63548.1	7705936 NT	AW027554.1	AW 503208.1	AI565065.1	AW965556.1	J00146.1	J00146.1	AF145712.1	U48724.1	AA527446.1		AA52/446.1	105,144,000.1	4504184 NT	AF003528.1	U64675.2	BE260201.1	A1278526.1	4504184 NT	4504184 NT	AB037747.1	AB037747.1	TN 8922968	8922965 NT		AB033104.1
Most Similar (Top) Hit BLAST E Value	1.0E-105	1.0E-105	1.0E-105	1.0E-105	1.0E-105	1.0E-105	1.0E-108	1.0E-106	1.0E-108	1.0E-106	1.0E-106	1.0E-106	1.0E-106	1.0E-106	10,	1.05-105	1.05-100	1.05-106	1.0E-106	1.0E-106	1.0E-108	1.0E-108	1.0E-108	1.0E-106	1.0E-106	1.0E-106	1.0E-106	1.0E-106	1.0E-106	1.0E-106
Expression Signal	1.8	2.89	5.44	1.8	2.06	2.01	0.98	1.54	1.77	0.79	1.21	2.66	4.51	5.12	0,0	9.12	8	8.39	1.63	1.26	1.94	4.23	2.97	2.97	5.01	5.01	2.38	2.36	0.8	0.98
ORF SEQ ID NO:	26987	27304	28428		28740	28968		19987	20286	20334	20334	21272	21444	21533	7,0970	21034	2007	22052	22229	22322	22324	22487	21183	21184	22635	22636	22867	22868	23053	23109
Exan SEQ ID NO:	16773	17111	18181	18430	18469	18679	L	10170	L		10527		11576	11661		1001	ł	12103	12333	12429	ŀ			11319	12837	12837	13068	13068	13248	13311
Probe SEQ ID NO:	6894	7234	8303	8560	8602	8867	145	198	531	689	9	1508	1874	1762	4700	2075	2 2	3	2456	2557	2559	2729	27.05	2795	2911	2911	3143	3143	3328	3394

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
3384		23110	0.98	1.0E-106	AB033104.1	N-	Homo sapiens mRNA for KIAA1278 protein, pertial cds
3952		23634	7.22	1.0E-106	AW974650.1	EST_HUMAN	EST386875 MAGE resequences, MAGN Homo saplens cDNA
3952	13860	23635	7.22	1.0E-108	AW974650.1	EST_HUMAN	EST386875 MAGE resequences, MAGN Homo sapiens cDNA
4497	L	24176	1.21	1.0E-106	BE144286.1	EST_HUMAN	MR0-HT0165-140200-008-d10 HT0165 Home sapiens cDNA
5183	15047		1.21	1.0E-106	L41644.1	NT	Homo sapiens dystrophin gene, exon 41
5298	15219	25022	2.98	1.0E-108	AA781155.1	EST HUMAN	aj24b09.s1 Soares, testis_NHT Homo sapiens cDNA clone 1391225.3' similar to gb:X12433 PROTEIN PHPS1-2 (HUMAN);
5711	L		6.78	1.0E-108		EST HUMAN	602154012F1 NIH_MGC_83 Homo sapiens cDNA done IMAGE:4285067 5
5866	15772		16.4	1.0E-108	11545913 NT	TN	Homo sapiens xylosyltransferase II (XT2), mRNA
5868	L		16.4	1.0E-108	11545913 NT	ĮŽ	Homo sapiens xylosytransferase II (XT2), mRNA
6361	16224	26385	5.59	1.0E-106	AA663779.1	EST HUMAN	ae72e07.s1 Stratagene schizo brain S11 Homo sapiens cDNA clone IMAGE:969732.3' similar to gb:X85873 KINESIN HEAVY CHAIN (HUMAN);
6390	16252	26412	4.83	1.0E-106	11429617 NT	LN	Homo sapiens XPMC2 protein (LOC57109), mRNA
6431	16292	26453	1.35	1.0E-106	BE292	EST_HUMAN	601105736F1 NIH_MGC_15 Homo sapiens cDNA done IMAGE:2988345 5'
6490		26517	7.6	1.0E-106	11425503 NT	K	Homo sapiens sorting nextn 11 (SNX11), mRNA
6490	16348	26518	7.6	1.0E-106	11425503 NT	NT	Homo saplens sorting nexin 11 (SNX11), mRNA
6657	.16537	26733	5.33	1.0E-108	BE741408.1	EST_HUMAN	601594331F1 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:3948463 5'
6657	16537	26734	5.33	1.0E-108	BE741408.1	EST_HUMAN	601594331F1 NIH_MGC_9 Homo sapiens cDNA done IMAGE:3948463 5'
6754	16633	26821	1.48	1.0E-108	AJ523068.1	EST_HUMAN	ar68a07.x1 Barstead aorta HPLRB6 Homo sabiens cDNA clone IMAGE:2127732 3' similar to gb:X06233 CALGRANULIN B (HUMAN);
7052	16929	27120	3.16	1.0E-106	Al654123.1	EST HUMAN	ty62a05.x1 NCI_CGAP_Kid11 Homo sapiens/cDNA clone IMAGE:2283832 3' similar to SW:ICA6_HUMAN_ Q05084 69 KD ISLET CELL AUTOANTIGEN
7281	L	27353	1.86	1.0E-106 /	AA825307.1	EST HUMAN	oc87e08.s1 NCI_CGAP_GCB1 Homo saplens cDNA clone IMAGE:1354790 3'
7281	17168	27354	1.86	1.0E-108	AA825307.1	EST_HUMAN	cc67e08.s1 NCI_CGAP_GCB1 Homo saplens cDNA clone IMAGE:1354790 3'
7361	17219	27419	2.79	1.0E-106	AI750447.1	EST_HUMAN	cr03a04.y1 Normal Human Trabecular Bone Cells Homo sapiens cDNA clone NHTBC_cn03a04 random
7424	17291	27501	1.86	1.0E-108	AI479569.1	EST_HUMAN	the 1102.x1 NC_CGAP_KId11 Homo sepiens cDNA clone IMAGE:2160699 3' similar to contains MSR1.t3 TAR1 PTR5 repetitive element;
7424	17291	27502	1.86	1.0E-106	1.0E-106 AI478569.1	EST HUMAN	tm41f02.x1 NCI_CGAP_Kid11 Homo sepiens cDNA clone IMAGE:2160699 3' similar to contains MSR1.t3 TAR1 PTR5 repetitive element:
7803	ı	27890	1.32	1.0E-108	BF027310.1	EST HUMAN	601671674F1 NIH_MGC_20 Homo sapiens cDNA clone IMAGE:3954403 5'
7803		27891	1.32	1.0E-106	BF027310.1	EST_HUMAN	601671674F1 NIH_MGC_20 Homo Saplens cDNA clane IMAGE:3954403 5'
7880		27975	5.83	1.0E-106	_	EST_HUMAN	np57b10.s1 NCL_CGAP_Br2 Homo sapiens cDNA clone IMAGE:1130395 3'
7880	17730	27976	5.83	1.0E-106	AA604417.1	EST_HUMAN	np57b10.s1 NCI_CGAP_Br2 Homo sapiens cDNA clone IMAGE:1130395 3'

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Table 4
Single Exon Probes Expressed In Heart

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	Top Hit Descriptor	Ig10d06.x1 NCI_CGAP_CLL1 Homo sapiens cDNA clone IMAGE:2108363 3' similar to SW:AACT_DICDI P05095 ALPHA-ACTININ 3, NON MUSCULAIX;	Homo saplens neuroendocrine-specific protein (NSP) gene, exon 4	602123963F1 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:4281039 5'	601066681F1 NIH_MGC_10 Homo saplens cDNA clone IMAGE:3452829 5	Homo sapiens HSPC049 protein (HSPC049), inRNA	Homo sapiens HSPC049 protein (HSPC049), mRNA	ze45e01.s1 Soares retina N2b4HR Homo sapiens cDNA clone IMAGE:361944 3' similar to contains THR.b1	THR repetitive element;	601177018F1 NIH_MGC_17 Homo sapiens cIDNA clone IMAGE:3532348 5	Homo sapiens NF2 gene	(1871-10 X1 NCI_CGAP_Pr28 Homo sepiens c) NA clone IMAGE:2248938 3' similar to gb:M14219 BONE PROTEOGLYCAN II PRECURSOR (HUMAN);	tt91e10x1 NCI_CGAP_Pr28 Homo sapiens cDNA clone IMAGE:2248938 3' similar to gb:M14219 BONE	PROTEOGLYCAN II PRECURSOR (HUMAN);	bb25b10.x1 NIH_MGC_14 Homo sapiens cDNA clone IMAGE:2963899 3' similar to gb:X53777 60S RIBOSOMAL PROTEIN L23 (HUMAN); gb:J05277 Mouse hexokinase mRNA, complete cds (MOUSE);	Homo saplens potassium channel subunit (HERG-3) mRNA, complete cds	Homo sapiens potassium channel subunit (HERG-3) mRNA, complete cds	Homo saplens pericentriolar material 1 (PCM1) mRNA	hi12a11.x1 NCI_CGAP_GU1 Homo sapiens cDNA clone IMAGE:2972060 3' similar to SW:3BP1_MOUSE P65194 SH3-BINDING PROTEIN 3BP-1.;	Human hepatocyte nuclear factor 4-alpha gene, exon 2	Human hepatocyte nuclear factor 4-alpha gene, exon 2	Homo sapiens KIAA0187 gene product (KIAA0187), mRNA	Horno sapiens PSN1 gene, alternative transcript	RC0-HT0372-241189-031-d03 HT0372 Homo saplens cDNA	801444922F1 NIH_MGC_65 Homo sapiens cDNA clone IMAGE:3848980 5'	601444922F1 NIH_MGC_65 Homo sapiens cDNA done IMAGE:3848980 51	Homo sapiens FYVE domain-containing dual specificity protein phosphatase FYVE-DSP2 mRNA, complete ods	Homo saplens FYVE domain-containing dual specificity protein phosphatase FYVE-DSP2 mRNA, complete cds
0	Top Hit Database Source	EST_HUMAN	TN	EST_HUMAN	EST_HUMAN	NT	NT		EST_HUMAN	EST_HUMAN	IN	EST HUMAN	-	EST_HUMAN	EST HUMAN	LN	NT TN	٦N	EST_HUMAN	FZ	5	Ę	Ę	EST_HUMAN	EST_HUMAN	EST_HUMAN	Þ	Ę
	Top Hit Acession No.	Al392850.1	L49141.1	BF666511.1	BE540550.1	11419701 NT	11419701 NT			BE296042.1	Y18000.1	AI886040.1		AI686040.1	BE206694.1	AF032897.1	AF032897.1	5453855 NT	AW664438.1	U72961.1	U72981.1	T661979 NT	AJ008005.1	AW384094.1	BE869016.1	BE869016.1	AF264717.1	AF264717.1
	Most Similar (Top) Hit BLAST E Value	1.0E-107	1.0E-107	1.0E-107		1.0E-107	1.0E-107			1.0E-108	1.0E-108	1.0E-108		1.0E-108	1.0E-108			1.0E-108		1.0E-108		1.0E-108			1.0E-108	1.0E-108	1.0E-108	1.0E-108
	Expression Signal	2.73	1.82	1.98	9.12	4.21	4.21		3.94	1.48	1.55	7.41		7.41		480	16.0	0.92	1.33	1.89	1.89	2.85	2.19	1.53	277	2.77	5.08	5.06
	ORF SEQ ID NO:	28228					28123			20711	20889	22063	l	22064	22150					24108							25745	
	Exen SEQ ID NO:	17978	18193	ı	L		17881		19873	10864	11151	12168	1	12166	12258	1		ļ	J.,	14320	14320		L	l_	L		15840	1
	Probe SEQ ID NO:	8087	8318	8327	8638	7698	8697		9187	626	1244	2282		2282	2378	3305	3305	3742	4065	4425	4425	4699	4868	5366	5393	5393	5732	6732

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Single Exon Probes Expressed in Heart

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	Top Hit Descriptor	Homo sapiens caveolin-1/-2 locus, Contig1, D75522, genes CAV2 (exons 1, 2a, and 2b), CAV1 (exons 1 and 2)	Homo sapiens G protein-coupled receptor, family C, group 5, member B (GPRC5B), mRNA	Homo sapiens delta-8 fatty acid desaturase (FADSD8) mRNA	Homo sapiens connective tissue growth factor-like protein precursor, mRNA, complete cds	Homo sapiens mRNA for Golgi-associated microtubule-binding protein (GMAP-210)	EST378258 MAGE resequences, MAGI Homo sapiens cDNA	Homo sapiens G protein-coupled receptor 48 (GPR48), mRNA	ttg1e10.x1 NCI_CGAP_Pr28 Homo eapliens cDNA clone IMAGE:2248938 3' similar to gb:M14219 BONE PROTEOCLYCAN II PRECURSOR (HUMAN):	#91e10.x1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:2248938 3' similar to gb:M14219 BONE	TACLECCE TOWN IF THE CONTROL OF THE	Home sapiens michal for PLUUUU3/ protein, pertea cds	602018571F1 NCL CGAP Brid? Home saplens cDNA clone IMAGE:4154297 5	Human mRNA for KIAA0220 gene, partial cds	Homo saplens hypothetical protein FLJ11316 (FLJ11316), mRNA	Homo sapiens reticulocalbin 1, EF-hand calcium binding domain (RCN1), mRNA	Homo saplens tetratricopeptide repeat domain 2 (TTC2) mRNA	Homo sapiens mRNA for KIAA0999 protein, partial cds	Homo sapiens mRNA for KIAA0999 protein, partial cds	Homo sapiens nucleolar phosphoprotein B23 (NPM1) mRNA, complete cds	Homo sapiens nucleolar phosphoprotein B23 (NPM1) mRNA, complete cds	Homo sapiens mRNA for KIAA0018 protein, partial cds	Homo sapiens chromosome 21 segment HS21 C084	Homo sapiens SNF5/IN1 gene, exon 8	ow95a01.x1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:1054536 3' similar to TR:002197 C02197 CIRCULATING CATHODIC ANTIGEN.	ow95a01.x1 Soares_fetal_liver_spleen_1NFLS;_S1 Homo sapiens cDNA clone IMAGE:1654536 3' similar to TR:002197 C02197 CIRCULATING CATHODIC ANTIGEN.	Homo sapiens guanylate cyclase activator 1A (retina) (GUCA1A) mRNA	J2816F Human fetal heart, Lambda ZAP Express Homo sapiens cDNA done J2816 5' similar to ZINC FINGER PROTEIN 2NEA.	CM3-NN0009-190400-150-110 NN0009 Homo seniens CDNA	CM3-NN0009-190400-150-f10 NN0009 Hamp sapiens cDNA	
Ē.	Database Source	FZ	NT	NT	N	N	EST HUMAN	1	EST HUMAN	MANUL FOR	-'I.	- N	EST_HUMAN	LΝ	LN L	NT .	TN	TN	LΝ	LN	LN	NT	LN	TN	EST_HUMAN	EST HUMAN	LN	EST LIMAN	EST HUMAN	EST_HUMAN	
	Top Hit Acession No.	AJ133269.1	11431857 NT	4758333 NT	AF083500.1	Y12490.1	4W966185.1	11441465 NT	A 1686040.1	Negeon 4	41000040.1	4KUZ4447.1	3F346356.1	386974.1	11422486 NT	11438391 NT	4507712 NT	4B023216.1	AB023216.1	A28699.1	A28699.1	013643.2	NL163284.2	(17123.1	A1022328.1	A1022328.1	4504206 NT	185100 4	1.0E-109 AW893192.1	1.0E-109 AW893192.1	
Most Similar	(Top) Hit BLAST E Value	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	100	1.05-100	1.0E-108	1.0E-108	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109		1 05-400	1.0E-109	1.0E-109	
	Expression Signal	1.37	5.35	3.34	1.93	2.77	4.26	2.03	4.14	77.7	2 20	2.18	9.58	6.0	0.92	1.51	3.64	14.64	14.64	9.63	4.89	1.48	2.03	1.87	3.88	3.88	2.76	88.1	1.46	1.45	
	ORF SEQ ID NO:	25804	26265	26421		24891	28731		22063	22084	25000	20003		19856	١		20219	20328	20329	20941	20941	21602	21985	21993	22344	22345	22346	22740	23068	23069	
Exa	0)	15695	16113	16280	16578	15099	18461	18565	12168	12168	4000	100770	0/281	10044	10183	10192				11095	11095	11728	12081	12091	12462	12452	12453	12048	13262	11	
Probe	SEQ ID NO:	5789	6247	6388	8698	8202	8533	8677	8706	8708	2000	8520 0738	08/A	88	212	22	459	583	883	1184	1185	1831	2194	2204	2681	2581	2682	υσυε	3342	3342	:

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Top Hit Descriptor	Homo sapiens retinol dehydrogenase homolog įsoform-1 (RDH) mRNA, complete cds	MR0-HT0209-110400-108-a04 HT0209 Homo sapiens cDNA	Homo saplens mRNA for KIAA0609 protein, partial cds	Homo sapiens mRNA for KIAA0609 protein, pertial cds	ts98e06.x1 NCI_CGAP_GC6 Homo sapiens cDNA clone IMAGE:2239330 3' similar to WP:F53A2.8	CEIOIUU.	nu93c12.s1 NCI_CGAP_Pr22 Homo sepiens ¿DNA clone IMAGE:1218262 3' similar to SW:GTT2_HUMAN P30712 GLUTHATHIONE S-TRANSFERASE THETA 2 ;	nu93c12.s1 NCI_CGAP_Pr22 Homo sapiens cDNA clone IMAGE:1218262 3' similar to SW:GTT2_HUMAN P30712 GLUTHATHIONE S-TRANSFERASE THETA 2;	Homo sapiens guanylate cyclase activator 1A (retina) (GUCA1A) mRNA	Homo sapiens KIAA0377 gene product (KIAA0377), mRNA	ya48e06.r1 Soares Infant brain 1NIB Homo sapiens cDNA clone IMAGE:53057 5'	601186922F2 NIH_MGC_15 Homo saplens cDNA clone IMAGE:2859636 6'	601186922F2 NIH_MGC_15 Homo septens cDNA clone IMAGE:2959636 5'	Homo sapiens placental protein 11 (serine proteinase) (P11) mRNA	RC1-HT0615-200400-022-d04 HT0615 Homo saplens cDNA	Homo saplens AT-binding transcription factor (1 (ATBF1), mRNA	601809495F1 NIH_MGC_18 Homo saplens cDNA clone IMAGE:4040279 6'	601809495F1 NIH_MGC_18 Homo sapiens cDNA clone IMAGE:4040279 5'	Noval human gene mapping to chomosome 13	PM0-BT0340-091299-002-e05 BT0340 Homo sapiens cDNA	7B18H01 Chromosome 7 Fetal Brain cDNA Library Homo sapiens cDNA clone 7B18H01	601479417F1 NIH_MGC_68 Homo sapiens cDNA clone IMAGE:3882124 6'	601479417F1 NIH_MGC_68 Homo sapiens cDNA clone IMAGE:3882124 5"	ys90g08.r1 Soares retina N2b5HR Homo sapiens cDNA clone IMAGE:222110 5' similar to SP:A53491 A 54401 RI IMETANIDE SENSITIVE NA K.C.1 COTRANSPORTER SOUNY	HSC1EC121 normalized infant brain cDNA Homo sepiens cDNA clone c-1ec12	601063030F1 NIH_MGC_10 Homo saplens cDNA clone IMAGE:3449599 5'	601063030F1 NIH_MGC_10 Homo sapiens cDNA clone IMAGE:3449599 5'	802080724F2 NIH_MGC_81 Homo sapiens ¿DNA done IMAGE:4245341 5'	Homo sapiens KIAA0744 gene product; histone deacetylase 7 (KIAA0744), mRNA	Homo sapiens KIAA0744 gene producț histone deacetylase 7 (KIAA0744), mRNA	AU121370 HEMBB1 Homo saplens cDNA clone HEMBB1002890 5'
Top Hit Database Source	Ę	EST_HUMAN	Г	Z E		ESI HUMAN	EST_HUMAN	EST HUMAN			EST_HUMAN	EST_HUMAN	EST_HUMAN		HUMAN		HUMAN				EST_HUMAN				EST HUMAN	7	Г	EST_HUMAN			EST_HUMAN
Top Hit Acession No.	4F240698.1	3E146144.1	4B011181.2	AB011181.2		41655417.1	4A662274.1	4A662274.1	4504208 NT	7662083 NT	315400.1	3E293673.1	BE293673.1	5174622 NT	3E179356.1	11432574 NT	3F182707.1	3F182707.1	4L049784.1	AW749130.1	AA077498.1	3E787540.1	3E787540.1	Lozopa 4		1.6	BE540909.1	3F694831.1	7662279 NT	7662279 NT	AU121370.1
Most Similar (Top) Hit BLAST E Value	1.0E-109	1.0E-109	1.0E-109	1.0E-109	10.4	1.05-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1 05 400	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109
Expression Signal	1.2	1.53	1.54	1.54	6	3.6/	1.02	1.02	2.25	1.19	1.04	98'0	0.88	2.31	1.48	3.66	5.01	5.01	1.36	1.23	1.72	5.71	5.71	c	141	2.93	2.93	14.2	2.12	2.12	1.88
ORF SEQ ID NO:	23195		23601	23602	100	23/32	23749	23750	23986	24175	24503	24626	24627	24950		26491	26492	26493	26820	26892		27139	27140	00770		28294	28295	28318	28464	28465	28606
Exan SEQ ID NO:	13390	13680	L	ł	1	13830	13972	13972	14202	14380	14720	14859	14859	15176	15364	16325	16326	16326	16632	16699	16907	16948	16948	47000	1	1_	L	18069	18212	18212	18341
Probe SEQ ID NO:	3474	3767	3911	3911	1	4034	4070	4070	4304	4486	4839	4984	4984	5254	5648	6465	6466	9949	6753	6820	2030	7071	1071	20.07	7482	8155	8155	8183	8335	8335	8468

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Top Hit Descriptor	Homo sapiens Chediak-Higashi syndrome 1 (CHS1) mRNA		Homo sapiens SNF5/IN11 gene, exon 6	Homo saplens gene for AF-6, complete cds	Homo sapiens defodinase, todothyronine, type II (DIO2), transcript variant 2, mRNA	Homo sapiens leucine-zipper-like transcriptional regulator, 1 (LZTR1), mRNA			Homo sapiens deiodinase, iodothyronine, type IJ (DIO2), transcript variant 2, mRNA	Human mRNA for Inward rectifier potassium channel, complete cds	Human dystrobrevin (DTN) gene, exon 20	Homo sapiens calcitonin receptor-like (CALCRL) mRNA	Homo saplens BAZ1B mRNA for bromodomain adjacent to zinc finger domain 1B, complete cds				Homo saplens Bruton's tyrosine kinase (BTK), alpha-D-galactosidase A (GLA), L44-like ribosomal protein	(L44L) and F1P3 (F1P3) genes, complete cds	Homo sapiens pregnancy-zone protein (PZP), mRNA	Homo sapiens pregnancy-zone protein (PZP), mRNA	bb82a05.y1 NIH_MGC_10 Hamo sapiens cDNA clone IMAGE:3048848 5' similar to TR:060312 060312 KIAA0568 PROTEIN;		Г	Г	Г	Homo sapiens hypothetical protein FLJ10300 (FLJ10300), mRNA	Homo saplens hypothetical protein FLJ10300 (FLJ10300), mRNA	Human cystic fibrosis transmembrane conductance regulator (CFTR) gene, exon 7		П	Homo sepiens mRNA for KIAA0868 protein, partial cds
Top Hit Database Source	NT	EST_HUMAN	TN	N	TN	TN		EST_HUMAN	LN	ΤN	TN	FZ	LΖ	EST_HUMAN	EST_HUMAN	LN	. !	Ł	NT	NT	EST HUMAN	EST HUMAN	EST HUMAN	N.	EST_HUMAN	N	NT	TN	EST_HUMAN	EST_HUMAN	<u>F</u>
Top Hit Acession No.	4502838 NT	W16510.1	Y17123.1	1.0E-109 AB011399.1	7549804 NT	5803073 NT	5803073 NT	C04498.1	7549804 NT	D87291.1	U84550.1	5031620 NT	1.0E-110 AB032253.1	1.0E-110 BE379477.1	BF508898.1	4503098 NT		8027	11436041 NT	11436041 NT	1.0E-110 BE018556.1	Al017213.1	1.0E-110 AU117812.1	7662441 NT	1.0E-110 BE299406.1	11419323 NT	11419323 NT	1.0E-110 M55112.1	AV714276.1	1.0E-110 AV714276.1	AB020675.1
Most Similar (Top) Hit BLAST E Value	1.0E-109	1.0E-109 W	1.0E-109 Y1	1.0E-109	1.0E-110	1.0E-110	1.0E-110	1.0E-110	1.0E-110	1.0E-110 D87291.1	1.0E-110 U84550.1	1.0E-110	1.0E-110	1.0E-110	1,0E-110 BF	1.0E-110		1.0E-110 U7	1.0E-110	1.0E-110	1.0E-110	1.0E-110 A	1.0E-110	1.0E-110	1.0E-110	1.0E-110	1.0E-110	1.0E-110	1.0E-110		1.0E-110
Expression Signal	2,19	4.83	1.65	2.8	1.48	3.88	3.88	30.63	1.67	16.0	1.16	- 8:	1.01	1.5	1.86	1.02		1.07	1.87	1.87	0.92	2.08	2.08	234	1.9	7.34	7.34	4.36	10.04	10.04	2.7
ORF SEQ ID NO:	28848		21993	25276	19781	19817	19818	19876	19781	20076															24929		_		26413		26432
Exan SEQ ID NO:	18564		1	L	0666	10021	10021	10059	0666	10255			11165		L	L	١		13083	13083	l	İ		L	L	L	上		L	16253	18270
Probe SEQ ID NO:	8676	8710	9259	9694	8	34	32	75	104	291	516	1163	1258	1879	2012	2810		3048	3158	3158	3966	4533	4555		5237	5511	5511	6032	6391	6391	6408

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Top Hit Descriptor	QV2-LT0053-020400-119-e04 LT0053 Homo sapiens cDNA	Homo sapiens galactokinase 2 (GALK2), mRNA	H.sapiens mRNA for myotonic dystrophy protein kinase like protein	601565604F1 NIH_MGC_21 Homo sepiens cDNA clone IMAGE:3840433 5'	601565604F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3840433 5	Zw67g02.r/ Soares_tests_NHT Homo sapiens cDNA clone IMAGE:781298 5' similar to TR:G1145816	601430784F1 NIH MGC 72 Homo seplens cPMA clone IMAGE 3024548 51	II.0-BT0163-04089-094-010 BT0163 Homo saniens cDNA	Homo saplens gene for AF-6, complete cds	PM3-NN1082-140900-006-f12 NN1082 Homo sapiens cDNA	Human ribosomal protein L23a mRNA, complete cds	Homo saplens ras GTPase activating protein-II(se (NGAP) mRNA	601458531F1 NIH_MGC_66 Homo sapiens cDNA clone IMAGE:3862086 5'	Homo saplens cat eye syndrome critical region gene 1 (CECR1), mRNA	Human cardiac alpha-myosin heavy chain (MYH6) gene, exons 32 to 34	Homo sapiens sex comb on midleg homolog 1 (SCMHI), mRNA	Homo saplens sex comb on midleg homolog 1 (SCMH1); mRNA	Homo sapiens DKFZP434D156 protein (DKFZP434D156), mRNA	Human enkephalin B (enkB) gene, exon 4 and 3' flank'and complete cds	247b07.rt Soares_pregnant_uterus_NbHPU Homo sablens cDNA done IMAGE:505045 5' similar to gb:M23575 PREGNANCY-SPECIFIC BETA-1 GLYCOPROTEIN C PRECURSOR (HUMAN);	247b07.rl Soares_pregnant_uterus_NbHPU Homo sapiens cDNA clone IMAGE:505045 5' similar to gb:M23575 PREGNANCY-SPECIFIC BETA-! GLYCOPROTEIN C PRECURSOR (HUMAN);	qp09g12.X1 NCI_CGAP_Kid5 Homo sapiens :DNA clone IMAGE:1917574 3' similar to gb:M29893 RAS- RELATED PROTEIN RALA (HUMAN):	ILZ-NT0101-280700-114-E03 NT0101 Homo sapiens cDNA	zn62c12.r1 Stratagene muscle 937209 Homo; saplens cDNA clone IMAGE:562774 5' similær to gb:X03740 MYOSIN HEAVY CHAIN, SKELETAL MUSCLE (HUMAN);	Human beta4-Integrin (ITGB4) gene, exon 13	601847132F1 NIH_MGC_55 Homo sapiens cDNA clone IMAGE:4078303 5	Human mRNA for integrin alpha-2 subunit	Human mRNA for integrin alpha-2 subunit	Homo sapiens Trio isoform mRNA, complete cds
Top Hit Database Source	EST_HUMAN	F	Ę	EST HUMAN	EST_HUMAN	1	EST HUMAN	EST HIMAN	N L	EST HUMAN	N FN		EST_HUMAN	N L	Ę	N F	, LZ	LN T	Z	EST HUMAN	EST HUMAN	HST HIMAN	EST HUMAN	EST HUMAN	LN	EST_HUMAN	TA	NT	TN
Top Hit Acession No.	1.0E-110 AW838394.1	11432732 NT	/12337.1	3E734357.1	1.0E-110 BE734357.1	7 00007	1.0E-110 AA440328.1	1.0E-110 DECS/210.1	1.0E-110 AB011399.1	1.0E-110 BF364546.1	J43701.1	4758807 NT	BF035327.1	8393092 NT	25142	6912641 NT	6912641 NT	7661569 NT	(02268.1	1.0E-111 AA151017.1	1.0E-111 AA151017.1	1344679 1	1.0E-111 BF368228.1			-	<17033.1		1.0E-111 AF091395.1
Most Similar (Top) Hit BLAST E	1.0E-110	1.0E-110	1.0E-110 Y12337.1	1.0E-110	1.0E-110	277	1.05-110	1 OF-110 /	1 OE-110 /	1.0E-110 E	1.0E-111 U	1.0E-111	1.0E-111	1.0E-111	1.0E-111 N	1.0E-111	1.0E-111	1.0E-111	1.0E-111 K02268.1	1.0E-111	1.0E-111	1 OF-441	1.0E-111	1.0E-111	1.0E-111 U66533.1	1.0E-111	1.0E-111 X17033.1	1.0E-111 X17033.1	1.0E-111
Expression Signal	2.91	4.27	3.7	3.49	3.49	9	2.43	87.8	1 83	5.07	16.85	1.02	1.87	3.58	73.62	1.17	1.17	1.08	4.45	2.82	2.82	1 74	3.03	2.29	3.13	10.8	13.75	13.75	3.26
ORF SEQ ID NO:	27683	28004	28266		28463		20102					19978		20489	20684	23342	23343	23760	23909	25117	25118	25885					27221		
Exen SEQ ID NO:	17374	ŀ	18018		18211	ł	17008		L	L	L	10161	10650	10658	10835	L	13556	13982	14133	15284	15284	15584	l	1	16674	16990	1_		17128
Probe SEQ ID NO:	7523	7915	8130	8334	8334	1 5	04.0	9204	9444	9578	168	189	718	728	911	3642	3642	4080	4235	5364	5364	5675	6402	6239	6795	7113	7149	7149	7251

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Top Hit Descriptor	as58g02.s1 NCI_CGAP_GCB1 Homo sapiens/cDNA clone IMACE:825170 3' similar to gb:L09235 VACUOLAR ATP SYNTHASE CATALYTIC SUBUNIT A, UBIQUITOUS (HUMAN);	2/3/101.r1 Scares_pregnant_uterus_NbHPU Homo saplens cDNA clone IMAGE:503545 5	Human thrombopoietin receptor (MPL) gene, exons 1,2,3,4,5 and 6	Homo saplens meningioma (disrupted in balanced translocation) 1 (MN1), mRNA	72C9 Human retina cDNA Tsp509I-cleaved sublibrary Homo saplens cDNA not directional	Homo sapiens low density lipoprotein-related protein 2 (LRP2), mRNA	Homo saptens low density lipoprotein-related protein 2 (LRP2), mRNA	Homo saplens acetyl-Coenzyme A carboxylase beta (ACACB), mRNA	Human steroidogenic acute regulatory protein (StAR) gene, exon 5	Human steroidogenio acute regulatory protein (StAR) gene, exon 5	UI-H-BI4-act-g-04-0-UI:s1 NCI_CGAP_Sub8 Homo sapiens cDNA clone IMAGE:3086023 3'	UI-H-BI4-act-g-04-0-UI.s1 NCI_CGAP_Sub8 Homo sapiens cDNA clone IMACE:3086023 3'	Homo sapiens HTRA serine protease (PRSS1/1) gene, complete cds	ZINC FINGER PROTEIN 135	Homo sapiens KIAA0440 protein (KIAA0440), mRNA	Homo sapiens KIAA0440 protein (KIAA0440), mRNA	801442874F1 NIH_MGC_65 Homo sapiens cDNA clone IMAGE:3846858 5	MR2-BT0590-090300-113-f09 BT0590 Homo sapiens cDNA	Homo sapiens mRNA for KIAA1411 protein, partial cds	Homo sapiens mRNA for KIAA1411 protein, partial cds	Homo sapiens low density lipoprotein receptor, related protein-deleted in tumor (LRPDIT), mRNA	ly35d07.r1 Soares melanocyte 2NbHM Homo sapiens cDNA clone IMAGE:273229 5	Homo sapiens solute carrier family 6 (neurotransmitter transporter, L-proline), member 7 (SLC6A7), mRNA	Homo sapiens solute carrier family 6 (neurotrensmitter transporter, L-proline), member 7 (SLC6A7), mRNA	AU118051 HEMBA1 Homo sapiens cDNA clone HEMBA1002773 5'	601443151F1 NIH_MGC_65 Hamo saplens cDNA clone IMAGE:3847285 5	601443161F1 NIH_MGC_65 Hamo sapiens cDNA clone IMAGE:3847285 5'	780g07.x1 Soares_NSF_F8_9W_OT_PA_P_S1 Homo sepiens cDNA clone IMAGE:3523020 3' similar to TR-Q9VW35 Q9VW35 CG8743 PROTEIN :	MR3-SN0009-100400-106-b12 SN0009 Horno sapiens cDNA	Homo sapiens mRNA for secreted modular calcium-binding protein (smoot gene)
Top Hit Database Source	EST_HUMAN	EST_HUMAN	LN TN	ΙN	EST_HUMAN	۲	LN.	Ę	TN	TN	EST_HUMAN	EST_HUMAN	F	SWISSPROT	뉟	Ę	EST_HUMAN	EST_HUMAN	뒫	NT	ĮŅ.	EST_HUMAN	FZ	Ę	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	EST HUMAN	NT
Top Hit Acession No.	1.0E-111 AA504160.1	.0E-111 AA131248.1	J68159.1	11417901 NT	W22562.1	11430460 NT	11430460 NT	4501854 NT			BF509039.1		1.0E-112 AF157623.1		7662125 NT	7662125 NT	BE866859.1	1.0E-112 BE076073.1	1.0E-112 AB037832.1	1.0E-112 AB037832.1	9055269 NT	N46046.1	11416777 NT	11416777 NT	1.0E-112 AU118051.1	BE867635.1	1.0E-112 BE867635.1	1.0E-112 BF111413.1	1.0E-112 AW863327.1	1.0E-112 AJ249900.1
Most Similar (Top) Hit BLAST E Value	1.0E-111	1.0E-111	1.0E-111 U68159.1	1.0E-111	1.0E-111 W	1.0E-111	1.0E-111	1.0E-112	1.0E-112 U29103.1	1.0E-112 U29103.1	1.0E-112 BI	1.0E-112	1.0E-112	1.0E-112 P52742	1.0E-112	1.0E-112	1.0E-112	1.0E-112	1.0E-112	1.0E-112	1.0E-112	1.0E-112 N	1.0E-112	1.0E-112	1.0E-112	1.0E-112	1.0E-112	1.0E-112	1.0E-112	1.0E-112
Expression Signal	1.51	6.35	4.52	3.04	1.69	1.39	1.39	2.66	4.49	4.49	1.48	1.48	2.78	1.72	5.88	5.88	2.45	0.83	5.12	5.12	0.94	33.34	1.81	1.81	1.65	2.25	2.25	2.09	4.25	2.86
ORF SEQ ID NO:	27935	27986	28529					L		L	20363					L		L	L	L			26364	26365						
Exan SEQ ID NO:	17690	17743	18277	18825		19422	1	1				L	L			L	_	L	L	١			16203	16203		1_	L	1	1	1 1
Probe SEQ ID NO:	7840	7893	8401	8038	9896	9959	8928	592	594	594	618	618	985	1046	1658	1658	2460	3809	4646	4646	5170	5477	6340	6340	6766	7181	7181	7893	8159	8318

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Probe SEQ ID NO: NO: 8445 8500 8500 8521 725 725 5035 5035 5035 5608 8550 8550 8550 8550 8550 8550 855	Exan SEQ ID NO: 18373 18373 10657 10657 10657 10657 10657 10657 10657 14907 14907 14907 15282 15282 15287 15287 15523 15	ORF SEQ ID NO: 28638 20487 20487 20487 20487 20487 20487 224677 24677 24677 24677 25813 25510 25510 25510 25510 25510 25510 25510 25510 25510 25510 25510 25510 25510 25510	Signal 1.75 1.75 1.75 1.76 1.29 1.29 1.29 1.29 1.29 1.29 1.29 1.29	Most Similar (Top) Hit BLAST E Value 1.0E-112 1.0E-112 1.0E-113 1.	Top Hit Acessian No. No. HE280479.1 A1792603.1 A1792603.1 A1792603.1 A1365586.1 A1365586.1 M11965.1 A1365586.1 A1365586.1 A1365586.1 A1365586.1 BF515218.1 A_J006976.1 A_J006976.1 A_J006976.1 A_J006976.1 A_J006976.1 A_J006976.1 A_J006976.1 A_J006976.1 BE382842.1 BE382842.1 BE382842.1 BE382842.1 BE382842.1 BE382842.1 BE382842.1 BE382842.1 BE382842.1 BE382842.1 BE382842.1 BE382842.1 BE382842.1 BE382842.1 BE382842.1	Top Hit Database Source Source T_HUMAN	Top Hit Descriptor 601155328F1 NIH_MGC_21 Homo sepiens cDNA clone IMAGE:189899 5' evideda, y5 NCI_CGAP_Kitd3 Homo sepiens cDNA clone IMAGE:1869902 5' similar to TR:C04362 Q64362 evideda, y5 NCI_CGAP_Kitd3 Homo sepiens cDNA clone IMAGE:1869902 5' similar to TR:C04362 Q64362 evideda, y5 NCI_CGAP_Kitd3 Homo sepiens cDNA clone IMAGE:1869902 5' similar to TR:C04362 Q64362 evideda, y5 NCI_CGAP_Kitd3 Homo sepiens cDNA clone IMAGE:1853625 3' evideda, y6 NCI_CGAP_Kitd3 Homo sepiens cDNA clone IMAGE:1953625 3' evideda, y6 NCI_CGAP_Kitd3 Homo sepiens cDNA clone IMAGE:1953625 3' evideda, y6 NCI_CGAP_Kitd3 Homo sepiens cDNA clone IMAGE:1953625 3' evideda, y6 NCI_CGAP_Kitd3 Homo sepiens cDNA clone IMAGE:1953625 3' evideda, y6 NCI_CGAP_Sid7 NCI_CGAP_Sid7 Homo sepiens cDNA clone IMAGE:1953625 3' evideda, y6 NCI_CGAP_Sid7 NCI_CGAP_Sid7 Homo sepiens cDNA clone IMAGE:1953625 3' evideda, y6 NCI_CGAP_Sid7 Homo sepiens cDNA clone IMAGE:195365 5' evideda, y6 NCI_CCAP_Sid7 Homo sepiens cDNA clone IMAGE:18736 5' evideda, y6 NCI_CCAP_Sid7 Homo sepiens cDNA clone IMAGE:18736 5' evideda, y6 NCI_CCAP_Sid7 Homo sepiens cDNA clone IMAGE:3872564 5' evideda, y6 NCI_C Homo sepiens cDNA clone IMAGE:3627564 5' evideda, y6 NCI_C Homo sepiens cDNA clone IMAGE:3627564 5' evideda, y6 NCI_C Homo sepiens cDNA clone IMAGE:3627564 5' evideda, y6 NCI_C Homo sepiens cDNA clone IMAGE:3627564 5' evideda, y6 NCI_C Homo sepiens cDNA clone IMAGE:3627564 5' evideda, y6 NCI_C Homo sepiens cDNA clone IMAGE:3627564 5' evideda, y6 NCI_C Homo sepiens cDNA clone IMAGE:3627564 5' evideda, y6 NCI_C Homo sepiens cDNA clone IMAGE:3627564 5' evideda, y6 NCI_C Homo sepiens cDNA clone IMAGE:3627564 5' evideda, y6 NCI_C Homo sepiens cDNA clone IMAGE:3627564 5' evideda, y6 NCI_C Homo sepiens cDNA clone IMAGE:3627564 5' evideda, y6 NCI_C Homo sepiens cDNA clone IMAGE:3627564 5' evideda, y6 NCI_C Homo sepiens cDNA clone IMAGE:3627564 5' evideda, y6 NCI_C Homo sepiens cDNA clone IMAGE:3627564 5' evideda, y6 NCI_C Homo sepiens cDNA clone IMAGE:36288366 6' evideda, y6 NCI_C Homo s
629	10588	20379	7.68	1.0E-114	T70551.1 ES	T HUMAN	yd15c01.s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:108288 3' similar to gb:A21187 ALPHA-2-MACROGLOBULIN PFECURSOR (HUMAN);contains Alu repetitive element; Homo saniens hypothetical protein FLJ20080 (FLJ20080), mRNA
1291 1648 2773				1.0E-114 1.0E-114	AB03310		Homo saplens in protretives protein in 252000 (1 1252000), in the saplens rhabdod tumor deletion region protein 1 (RTDR1), mRNA Homo saplens mRNA for KIAA1276 protein, partial cds

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Mo E B	.82 1.0E-114 AB033102.1 NT Homo saplens mRNA for KIAA1276 protein, partial cds		1.2 1.0E-114 BF206374.1 EST_HUMAN 601869932F1 NIH_MGC_19 Homo saplens cDNA clone IMAGE:4100214 5'	95 1.0E-114 AF149773.1 NT Homo sapiens NOD1 protein (NOD1) gene, exons 1, 2, and 3		1.0E-114 AF004849.1 NT		37 1.0E-114 4508880 NT (TM) and short cytoplasmic domain, (semaphorin) 54 (SEM45A) mRNA			08 1.0E-114 Y18000.1 NT Homo saplens NF2 gene	ı	1.0E-114 AI363139.1 EST_HUMAN	1.0E-114 Al363139.1		.35 1.0E-114 AB011133.1 NT Homo saplens mRNA for KIAA0561 protein, partial cds			1.0E-114 AF077754.1		ba73g12.yf·NIH_MGC_20 Homo saplens cDNA clone IMAGE:2906086 5' similar to gb:X17206 40S	RIBOSOMAL PROTEIN S4 (HUMAN); gb:Mx:0632 Mouse LLRep3 protein mRNA from a repetitive element, 14.0 pc:0440e200ses 1 lest uniman complete (MOUSE).	4 0E-114 AV733454 4 EST HIMAN	4 OF 444 AV7202454 4 FET III IMANI	1.0E-114 AV /33434.1 EGI HUMAN	1.0E-114 AV733454.1 EST_HUMAN	1.0E-114 AV733454.1 EST_HUMAN		1.0E-114 11034850 NT	.06 1.0E-114 11034850 NT Homo saplens hypothetical protein (DJ1042K 10.2), mRNA	1.0E-115 4758111 NT	.95 1.0E-118 4505938 NT Homo sapiens polymerase (RNA) II (DNA dirécted) polypeptide A (220kD) (POLR2A) mRNA
op Hit Acession No.								4506880	4506880			4557600		1									T	1				11418041	11034850	11034850	4758111	4505938
Most Similar (Top) Hit BLAST E Value	1.0E-114 A	1.0E-114 X	1.0E-114 B					1.0E-114	1.0E-114	۱>-	1.0E-114 Y	ı	l	1.0E-114 A	1.0E-114 U	1.0E-114 A	1.0E-114 A	1.0E-114 A	1.0E-114 A	1.0E-114 A		244	4 OE-144 A	A 544 A 5	A 4-11-4 A	1.0E-114 A	1.0E-114 A	1.0E-114	1.0E-114	1.0E-114	1.0E-115	1.0E-115
Expression Signal	0.82	2.29	1.2	1.95	12	2.31		1.37	1.37	7.08	7.08	1.86	1.73	1.73	3.39	6.35	6.35	3.79	3.13	1.31		7 4 7	4 58	3 7	8.9	2.86	2.86	3.21	3.06	3.06	3.36	0.95
ORF SEQ ID NO:	19824	22814	22859		24782			25040	25041		26339					27181				27940			98886				28933			25223		19920
Exon SEQ ID NO:		13019	13060	13843	14988	L	ł	15237	15237	16179	16179	16486	1	ı		16987	16967	17285	16459	17694		1004	18300	1000	1			19748				10099
Probe SEQ ID NO:	2773	3092	3135	3934	5120	5121		5316	5316	6316	6316	9089	6748	6748	7048	7090	7090	7418	7447	7844		0467	8527	2020	/700	8834	8834	8479	9729	9729	21	125

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Proba SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Detabase Source	Top Hit Descriptor
129	10103		1.99	1.0E-115	4557887	ΤN	Homo saplens keratin 18 (KRT18) mRNA
289	10253	20073	2.17	1.0E-115	AW804759.1	EST_HUMAN	QV4-UM0094-300300-156-b08 UM0094 Horn; sepiens cDNA
525	}		1.08	1.0E-115	A[339206.1	EST HUMAN	qt6f01.x1 NCI_CGAP_GC4 Home sapiens cDNA clone IMAGE:1946809 3' similar to TR:000536 000636 TTF-I INTERACTING PEPTIDE 5;
	1			1,77		TOTAL TOTAL	q06f01,x1 NCL_CGAP_GC4 Homo saplens cDNA clone IMAGE:1946809 3' similar to TR:000536 000536
929		202/B		1.05-115	AISSBZUG	NIWINIU I SE	I I I I I I I I I I I I I I I I I I I
2	1			1.0E-115		LN	Hamo capiens (Tanstorming grown ractor beta-activated tonase-binding protein (TAD), mixta
769	1			1.0E-115		N	Homo sapiens transforming growth factor beta-activated Kinase-binding protein 1 (1 AB1), mKWA
177	10701	20540	49.82	1.0E-115	4503794 NT	NT	Homo sapiens ferritin, heavy polypeptide 1 (FTiH1) mRNA
1539	11443	21301	0.92	1.0E-115	AF229180.1	IN	Homo sapiens alpha-aminoadipate semialdehyde synthase mRNA, complete cds
1539	11443	21302	0.92	1.0E-115	AF229180.1	NT	Homo sapiens alpha-aminoadipate semialdehyde synthase mRNA, complete cds
1785	11683	21561	3.14	1.0E-115	AJ277892.1	NT	Homo saplens partial TTN gene for titin
	1_						Homo sapiens Bruton's tyrosine kinase (BTK), alpha-D-galactosidase A (GLA), L44-like ribosomal protein
1798	11696	21573	1.42	1.0E-115	U78027.1	NT	(L44L) and FTP3 (FTP3) genes, complete cds
2034	11925		0.87	1.0E-115	BE745469.1	EST_HUMAN	601579838F1 NIH_MGC_9 Home sapiens cDNA clone IMAGE:3928832 5'
2034	11925	21818	0.87	1.0E-115	BE745469.1	EST_HUMAN	601579839F1 NIH_MGC_9 Homo saplens cDNA clone IMAGE:3928832 5'
2820	12749		1.78	1.0E-115	AW804759.1	EST_HUMAN	QV4-UM0094 300300-156-b08 UM0094 Homo septens cDNA
3077	13004	22795	2.1	1.0E-115	AJ245922.1	INT	Homo saplens mRNA for alpha-tubulin 8 (TUBA8 gene)
3077	13004	22796	2.1	1.0E-115	AJ245922.1	NT	Homo sapiens mRNA for alpha-tubulin 8 (TUBA8 gene)
3427	13344	23149	4.03	1.0E-115	AJ277892.1	N	Homo saplens partial TTN gene for titin
3958	١.	23640	4.04	1.0E-115		NT	Homo sapiens mRNA for KIAA0350 protein, tyartial cds
4169	1		1.08	1.0E-115	AL13716	NT	Novel human gene mapping to chomosome X
4301	14189		3.41	1.0E-115		۲N	Homo sapiens strZ-like 3 (SIRT3), mRNA
4335	14232	24014	3.83	1.0E-115	4758279 NT	TN	Homo saplens EphA4 (EPHA4) mRNA
4578	14468	24254		1.0E-115	AL096857.1	LN	Novel human mRNA from chromosome 1, which has similarities to BAT2 genes
4578	14468		2.58	1.0E-115	AL096857.1	IN	Novel human mRNA from chromosome 1, which has similarities to BAT2 genes
4813	14696			1.0E-115	AL163268.2	۲	Homo sapiens chromosome 21 segment HS21C068
4813	1.			1.0E-115		Ę	Homo sapiens chromosome 21 segment HS21C088
5279	L			1.0E-115	AW970335.1	EST_HUMAN	EST382416 MAGE resequences, MAGK Homo saplens cDNA
5330	L	L	7.22	1.0E-115	BF665387.1	EST_HUMAN	602119346F1 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:4276738 6'
5402	15321	25369	2.05	1.0E-115	11425128 NT	NT	Homo saplens similar to ER to nucleus signajling 1 (H. saplens) (LOC63433), mRNA
5402	15321	26370	2.05	1.0E-115	11425128 NT	٦	Homo sapiens similar to ER to nucleus signailing 1 (H. sapiens) (LOC63433), mRNA
5852	15758		12.92	1.0E-115	11426038 NT	Ę	Homo sapiens similar to ribosomal protein S26 (H. sapiens) (LOC63436), mRNA
6933	15838	25960	1.93	1.0E-115	7661883 NT	NT	Homo sapiens KIAA0054 gene product, Helicase (KIAA0054), mRNA

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	\neg	\neg	_			-			\neg									i	_	т		1					_			
Top Hit Descriptor	Homo sapiens KIAA0054 gene product, Helicase (KIAA0054), mRNA	oz31a06.x1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA clone IMAGE:1676914 3	oz31a06.x1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA clone IMAGE:1676914.3'	Homo sapiens mRNA for KIAA0995 protein, partial cds	RC6-ET0081-130700-011-G01 ET0081 Homo sapiens cDNA	RC6-ET0081-130700-011-G01 ET0081 Homo saplens cDNA	Homo sapiens eukaryotic translation initiation fector 4B (EIF4B), mRNA	Human mRNA for KIAA0338 gene, partial cds	Human mRNA for KIAA0338 gene, partial cds	xx32f08.x1 NCI_CGAP_Utf. Homo eapiens cDNA clone IMAGE:2839239 3' sImiler to SW:CAYP_CANFA P10463 CALCYPHOSINE';	Homo sapiens calcium channel, voltage-dependent, alpha 1E subunit (CACNA1E) mRNA	601121347F1 NIH_MGC_20 Homo sapiens cDNA done IMAGE:2988875 5'	Homo sapiens synaptojanin 1 (SYNJ1), mRNA	Homo saplens synaptojanin 1 (SYNJ1), mRNA	Homo sapians pericentrin (PCNT) mRNA	Homo saplens pertcentrin (PCNT) mRNA	AU133080 NT2RP4 Homo saplens cDNA clorie NT2RP4001228 5'	Human apolipoprotein B-100 (apoB) gene, exons 17 and 18	Human apolipoprotein B∙100 (apoB) gene, exons 17 and 18	Homo saplens protein phosphatase, EF hand calcium-binding domain 1 (PPEF1) mRNA	Human olfactory receptor offr17-201-1 (OR17-201-1) gene, olfactory receptor offr17-32 (OR17-32) gene and offectory receptor pseudo olfr17-01 (OR17-01) pseudogene, complete cds	Homo sapiens mRNA for KIAA0780 protein, partial cds	801513337F1 NIH_MGC_71 Homo sapiens cDNA done IMAGE:3914600 5	Homo sapiens DIGeorge syndrome critical region, centromeric end	Homo sapiens DiGeorge syndrome critical region, centromeric end	Homo sapiens sodium phosphate transporter 3 (NPT3) mRNA	PM-BT135-070499-016 BT135 Homo saplans cDNA	Homo sapiens partial 5-HT4 receptor gene, exons 2 to 5	zc24d07.r1 Soares_senescent_fibroblasts_NbHSF Homo sapiens cDNA clone IMAGE:323245 5' similar to sw:mDHM_MOUSE P08249 MALATE DEHYDROGENASE, MITOCHONDRIAL PRECURSOR;	Homo sapiens mRNA for KIAA1636 protein, partial cds
Top Hit Database Source	1. 1		EST_HUMAN	NT	EST HUMAN	EST_HUMAN	NT	TN	NT	EST HUMAN	N	EST_HUMAN	N-I	Z	LΝ	۲Z	EST_HUMAN	LN	TN	NT	Ę	Z	EST_HUMAN	Z	ΙZ	Z	EST HUMAN	Ŋ	EST_HUMAN	Ę
Top Hit Acession No.	7661883 NT	AI076598.1	AI076598.1	AB023212.1	BE830187.1	BE830187.1	11434772 NT	AB002336.1	AB002336.1	AW571544.1	4502528 NT	BE275502.1	4507334 NT	4507334 NT	5174478 NT	5174478 NT	AU133080.1	M19824.1	M19824.1	5453941 NT	1178308 1	AB018333.1	BE889256.1	L77570.1		5031954 NT	Al907096.1	AJ243213.1	W42822.1	AB046856.1
Most Similar (Top) Hit BLAST E Value	_		_		1.0E-115	1.0E-115	1.0E-115	1.0E-115	1.0E-116	1.0E-115				1.0E-116	1.0E-116	1.0E-116	1.0E-116	1.0E-116	1.0E-116	1.0E-116	1 0F-118			1.0E-116			1.0E-116	1.0E-116	1.0E-116	
Expression Signal	1.83	1.55	1.55	7.41	12.78	12.78	2.2	1.92	1.92	3.5	226	1.42	1.25	689	2.38	2.38	1.21	1.01	1.01	1.88	1 49	4.48	2.19	4.73	4.73	2.01		1.2	5.88	1.65
ORF SEQ ID NO:		26333	26334	26401	26811	26812	27184	27816	27817	28239]				21733		21765		21835	22042		22181					L		25641	
Exan SEQ ID NO:	Ш	16178	16176	16241	16623	16623	16993	17595	17595	17990	L	Ŀ	L		L	L	_	L		12143	<u> </u>	. _	L	L	13062	L	L	L	15550	1
Probe SEQ ID NO:	5933	6313	6313	6379	6744	6744	7116	7745	7745	8100	8701	560	783	841	1952	1952	1980	2050	2050	2259	2202	2407	2704	3137	3137	4281	4747	5122	5637	6788

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Тор Hit Descriptor	Homo sapiens mRNA for KIAA1636 protein, partial cds					EST62685 Jurkat T-cells V Homo saplens cDNA 5' end similar to similar to keratin 2	EST62685 Jurkat T-cells V Homo sapiens cDNA 5' end similar to similar to keratin 2	601338288F1 NIH_MGC_53 Homo saplens cDNA clone IMAGE:3680880 5	qh09c05.x1 Soeres_NFL_T_GBC_S1 Homo explens cDNA clone IMAGE:1844168 3' similar to gb.X53741_ma1 FIBULIN-1, ISOFORM A PRECURSOR (HUMAN);	Homo sapiens laminin, alpha 2 (merosin, congenital muscular dystrophy) (LAMA2), mRNA	CM2-CT0482-300800-349-e06 CT0482 Homo sepiens cDNA	qq41e04.x1 Soares_NhHMPu_S1 Homo saplens cDNA clone IMAGE:1935102 3' similar to WP:B0495.7 CE01765;	П	Г	Г	Mus musculus fragile-X-related protein 1 (Fx1h) gene, exons 13a through 15	Homo sapiens lymphocyte activation-associated protein mRNA, complete cds	Human apolipoprotein B-100 (apoB) gene, excin 10			EST188414 HCC cell line (matastasis to liver in mouse) II Homo sapiens cDNA 5' end similar to ribosomal	Т	7	1	TANDERS MINISTER OF THE PROPERTY OF THE PROPER	H.seplens mKNA for I PCK16 protein	Homo sapiens Scar2 (SCAR2) gene, partial cds	Homo saplens Scar2 (SCAR2) gene, partial cds	Homo sapiens mRNA for KIAA0866 protein, complete cds		Homo saplens nuclear hormone receptor (shp) gene, 3 end of cds	Homo sapiens nuclear hormone receptor (shp) gene, 3' end of cds
Top Hit Database Source	N	EST_HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	된	EST_HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	INT	TN	NT	NT	EST_HUMAN	EST_HUMAN	140731111 2502	SS. I ES I DOMONIA	EGT CITATAN	-1	z	NT	LN	NT	NT	EST_HUMAN	N	N
Top Hit Acession No.	AB046856.1	BF677910.1	BE158133.1	C02944.1	AV716314.1	AA354256.1	AA354256.1	BE565507.1	AI216352.1		3F3358	A1367140 1	. IW	AL134889.1	4826636 NT	AF124393.1	AF123320.1	M19816.1	AW957699.1	AA978114.1	, , , , , , , ,	AA3 107 23. 1	4 04440 4	X00070.1	X89670.1	X89670.1	AF134304.2	AF134304.2	AB020673.1	BE730508.1		L76571.1
Most Similar (Top) Hit BLAST E Value	1.0E-116 A	1.0E-116	1.0E-116	1.0E-116	1.0E-116	1.0E-116	1.0E-116	1.0E-116	1.0E-116	1.0E-116	1.0E-116	1 0F-116	1.0E-116	1.0E-116	l		1.0E-117	1.0E-117	1.0E-117	1.0E-117	277	1.0E-117	4 00 447			1.0E-117	1.0E-117	1.0E-117	1.0E-117	1.0E-117		1.0E-117
Expression Signal	1.65	72.79	1.8	3.59	76.7	1.99	1.99	1.43	1.98	1.77	3.68	3.33	2.08	268	1.18	1.69	6.25	2.3	2.99	1.75		3.02	00.1	1.80	1.35	1.35	9.22	9.22	3.36	2.5		4.99
ORF SEQ ID NO:	25782			26156							28218				20296	L										24300	24382			24978		28427
Exon SEQ ID NO:	15675		15829	16018	16139	l	L.	<u>l_</u>	L	L	L	L		L	L	L	l	L	L		L	\perp	┙			14510	14591		l	15202	Ш	16265
Probe SEQ ID NO:	5768	5853	5924	6145	6275	6874	6874	7173	7260	7577	8074	7777	9591	9776	546	1061	1719	1780	2164	3230		3908	474	44/5	4622	4622	4705	4705	4856	6280	6404	6404

WO 01/57274 PCT/US01/00666

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		Т	T		T	Т	Т	Т		Τ	Т	Т	Т	Τ	Т	Г	Т	Г	Τ	Ť.	Т	Ė	1	T	Ϊ	T	Ī	Ť	Ť	T	T	T	T
Onigio Exon Probas Expressed in near	Top Hit Descriptor	AV717788 DCB Homo sapiens cDNA clone DCBBAE01 5'	AV717788 DCB Homo saplens cDNA clone DCBBAE01 57	wp86b07 x1 NCI_CGAP_Brn25 Homo sapiens cDNA clone IMAGE:2468629 3' similar to TR:O75065 O75065 KIAA0477 PROTEIN.;	Homo sapiens neural cell adheston molecule 1 (NCAM1), mRNA	Homo sapiens neural cell adhesion molecule 1 (NCAM1), mRNA	Human gene for very low density lipoprotein reciptor, exon 11	601569317F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3843748 5'	zd83b11.r1 Soares_fetal_heart_NbHH19W Homo sepiens cDNA done IMAGE:347229 5' similar to gb:M14219 BONE PROTEOGLYCAN II PRECURSOR (HUMAN);	Homo sapiens mRNA for MEGF8, partial cds	Homo sapiens mRNA for MEGF8, partial cds	601186203F1 NIH_MGC_8 Homo sepiens cDNA clone IMAGE:3544296 5'	Homo saplens ATP-binding cassette, sub-family A (ABC1), member 3 (ABCA3), mRNA	Homo sapiens ATP-binding cassette, sub-family A (ABC1), member 3 (ABCA3), mRNA	Homo saplens HSPC151 mRNA, complete cds	DKFZp434I056_r1 434 (synonym: htes3) Homo saplens cDNA clone DKFZp434I056 5'	Homo sapiens hypothetical protein (DJ328E19 C1.1), mRNA	Homo sapiens sine oculis homeobox (Drosophila) homolog 1 (SIX1) mRNA	601281947F1 NIH_MGC_44 Homo saplens cDNA clone IMAGE:3604019 5'	601281947F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3604019 5'	601281947F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3604019 5'	EST363799 MAGE resequences, MAGB Homb sapiens cDNA	Human breakpoint cluster region (BCR) gene, complete cds	Human breakpoint cluster region (BCR) gene, complete cds	Homo sapiens PRKY exon 7	[qp01f05.x1 NC]_CGAP_Kid5 Homo sapiens cl2NA clone IMAGE:1916769 3'	ap01f05.x1 NCI_CGAP_Kid5 Homo sapiens cINA clone IMAGE:1916769 3'	Human mRNA for ribosomal protein, complete cds	Homo sapiens calcium channel gamma 4 subunit (CACNG4) gene, exon 3	Homo sapiens calcium channel gamma 4 subunit (CACNG4) gene, exon 3	Homo saplens transient receptor potential charinel 5 (TRPC5), mRNA	Homo saplens latent transforming growth factor beta binding protein 2 (LTBP2) mRNA	Homo sapiens latent transforming growth factor beta binding protein 2 (LTBP2) mRNA
שום באחזו ריום	Top Hit Database Source	EST_HUMAN	EST_HUMAN	EST HUMAN	1	NT	K	EST_HUMAN	EST_HUMAN	N	N FN	EST_HUMAN	NT	ΙΝ	NT	EST_HUMAN	NT	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	NT	TN	EST_HUMAN	EST_HUMAN	NT	NT	NT	LN	L	NT
	Top Hit Acession No.	AV717788.1	AV717788.1	A(950145.1	34989	10834989 NT	D16524.1	BE733922.1	W80605.1	AB011541.1	AB011541.1	BE269856.1	4501848 NT	4501848 NT	AF161500.1	AL045854.1	7657016 NT	74680	BE389705.1			AW951729.1		U07000.1	Y13932.1	Al347694.1	AI347694.1	D23660.1	AF142624.1	AF142624.1	11420764 NT	4857732 NT	4557732 NT
	Most Similar (Top) Hit BLAST E Value	1.0E-117	1.0E-117	1.0E-117	1.0E-117	1.0E-117	1.0E-117	1.0E-117	1.0E-117	1.0E-117	1.0E-117	1.0E-117	1.0E-117	1.0E-117	_	1.0E-118	1.0E-118		1.0E-118	_	1.0E-118	1.0E-118	1.0E-118	1.0E-118		_	1.0E-118	1.0E-118	1.0E-118	1.0E-118	1.0E-118	1.0E-118	1.0E-118
	Expression Signal	3.77	3.77	5.83	1.7	1.7	2.28	1.61	10.31	3.96	3.96	15.63	2.22	2.22	9.54	2.59	5.09	0.98	2.39	2.39	2.39	3.77	2.38	2.38	3.73	4.51	4.51	7.67	2.02	2.02	1.88	1.87	1.87
	ORF SEQ ID NO:	26482	26483	26725		26909	27428	27708	28512	28732	28733		28974	28975	19863	19890	20261	. 20667	21975	21976	21977		22465	22466		22887	22888	23679	25053	25054	25620	28054	26055
	Exan SEQ ID NO:	16316	16316	16531	16716		17228	17486	18262	18482	18462	18516					_						12574							15249	15535	15924	15924
	Probe SEQ ID NO:	6455	6455	6651	6837	6837	7361	7635	8385	8595	8595	8698	8872	8872	64	8	506	896	2186	2186	2186	2281	2711	2711	3066	3159	3159	3998	5329	5329	6620	6020	6020

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						-			4	4		Ir to SW:ZP3A_HUMAN	d similar to dynein, light								o WP:E04F6.2		-03					128 3' similar to			
Top Hit Descriptor	Homo sapiens chromosome 2 open reading fraine 3 (C2ORF3), mRNA	601469159F1 NIH_MGC_67 Hamo sepiens cDNA clane IMAGE:3872247 5	QV0-BT0263-090200-097-h03 BT0263 Homo sapiens cDNA	QV0-BT0263-080200-097-h03 BT0263 Homo sapiens cDNA	2x98d07.r1 Soares_NhHMPu_S1 Homo sapieris cDNA clone IMAGE:811789 5'	zx88d07.r1 Soares_NhHMPu_S1 Homo saplens cDNA clone IMAGE:811789 5'	Human mRNA for KIAA0383 gene, partial cds	Human mRNA for KIAA0383 gene, partial cds	Homo saplens latent transforming growth factor beta binding protein 2 (LTBP2) mRNA	Homo sapiens latent transforming growth facto; beta binding protein 2 (LTBP2) mRNA	601144863F2 NIH_MGC_19 Homo sapiens cDNA done IMAGE:3160502 5	7n17e09.x1 NCI_CGAP_Brn23 Homo sapiens CDNA clone IMAGE:3584785 3' similar to SW:ZP3A_HUMAN P21754 ZONA PELLUCIDA SPERM-BINDING PROTEIN 3A PRECURSOR;	EST186814 HCC cell line (matastasis to liver in mouse) Il Homo sapiens cDNA 5' end similar to dynein, light	chain 1, cytoplasmic	QV0-UM0091-120900-385-b12 UM0091 Homo sapiens cDNA	QV0-UM0091-120900-385-b12 UM0091 Homo saplens cDNA	Homo sapiens chloride channel CLC4 (ClC4) mRNA, complete cds	Homo sapiens CGI-105 protein (LOC51011), mRNA	Homo sapiens mRNA for KIAA0930 protein, partial cds	Homo sapiens hypothetical protein FLJ10052 (FLJ10052), mRNA	on10b05.s1 NCI_CGAP_Lu5 Homo saplens cDNA clone IMAGE:1556241 3' similar to WP:E04F6.2 CE01214;	Homo sapiens glutamate receptor, ionotropic, l'ainate 1 (GRIK1) mRNA	7B14F03 Chromosome 7 Felal Brain cDNA Library Homo sapiens cDNA clone 7B14F03	AU133399 NT2RP4 Homo sapiens cDNA clone NT2RP4001991 5'	Human neurofibromin (NF1) gene, complete ods	RC1-NN0073-250800-018-g08 NN0073 Homb sapiens cDNA	AV693731 GKC Homo sapiens cDNA clone GKCDHB03 5'	db77c09.x1 Soares fetal heart NbHH19W Homo septens cDNA clone IMAGE:1706128 3' similar to SW:K1CJ_MOUSE P02535 KERATIN, TYPE! I CYTOSKELETAL 10;	Human c-fes/fps proto-oncogene	EST386296 MAGE resequences, MAGM Horno sapiens cDNA	601592005F1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:3946081 5'
Top Hit Database Source	N	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	之	Ν	N	N	EST_HUMAN	EST HUMAN		EST_HUMAN	EST_HUMAN	EST_HUMAN	TN	LN	LN	N	30.1 EST HUMAN	Ϋ́	EST_HUMAN	EST_HUMAN	Z	EST HUMAN	EST HUMAN	EST_HUMAN	NT	EST_HUMAN	EST_HUMAN
Top Hit Acession No.	11431050 NT	BE781223.1	BE062855.1	BE062855.1	AA443024.1	AA443024.1	AB002381.1	AB002381.1	4557732 NT	4557732 NT	BE263134.1	BF195407.1		AA315007.1	BF093687.1	BF093687.1	AF170492.1	7705607 NT	AB023147.1	8922205 NT	AA916760.1	4504116	AA077394.1	AU133399.1	M89914.1	BE936121.1	AV693731.1	Al150703.1	X06292.1	AW974193.1	BE796614.1
Most Similar (Top) Hit BLAST E	1.0E-118	1.0E-118		1.0E-118	1.0E-118	1.0E-118	1.0E-118	_	1.0E-118	1.0E-118	1.0E-118	1.0E-118	•	_		1.0E-118	1.0E-119	1.0E-119	1.0E-119	1.0E-119	1.0E-119		1.0E-119	1.0E-119	1.0E-119	1.0E-119			1.0E-119	_	
Expression Signal	4.13	2.23	7.81	7.81	1.34	1.34	1.16	1.16	1.28	1.28	5.71	1.18		3.06	1.75	1.75	0.81	1.55	2.09	1.81	1.08	1.15	0.95	2.45	14.93	3.32	2.24	7.19	2.79	4.26	1.42
ORF SEQ ID NO:	26509			26961	26963	26964	27100	27101	27129	27130	27288	28027		28737	28947	28948	20508	20781	21663	22783		23573	24753	24969	24980	24984			25914	26925	26403
Exan SEQ ID NO:	16341	16528	16763	16763	16767	16767	16912		16939	16939	17100	17786		ļ	18659	18659	10672	12683	11786	12991	13126	13781	14979	15194	15204	15207	15256		L	ll	IJ
Probe SEQ ID NO:	6482	6648	6884	6884	6888	6888	7036	7035	7082	7062	7223	7936		8598	8847	8847	741	1021	1891	3064	3202	3870	5111	5272	5282	5285	5336	5726	5887	6895	6381

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rop Hit Descriptor	Human muscle glycogen phosphorylase (PYGM) gene, 5'UTR and exon 1	Homo saplens calcineurIn binding protein 1 (KIAA0330), mRNA	Homo sapiens NF2 gene	AU134963 PLACE1 Homo saplens cDNA clone PLACE1000899 6'	Homo sapiens TNF receptor-associated factor 1 (TRAF1) mRNA	Homo sapiens mRNA for KIAA0581 protein, partial cds	Homo sapiens inositol polyphosphate-4-phosphatase, type I, 107kD (INPP4A), splice variant a, mRNA	Homo sapiens inositol polyphosphate-4-phosphatase, type I, 107kD (INPP4A), splice variant a, mRNA	Homo sapiens metabotropic glutamate receptor 1 beta (mGluR1beta) mRNA, complete cds	602014759F1 NCI_CGAP_Brn64 Homo sapiens cDNA clone IMAGE:4150286 5	602014759F1 NCI_CGAP_Brn64 Homo sapiens cDNA clone IMAGE:4150286 5'	Homo sapiens hHb3 gene for hair keratin, exons 1 to 9	Homo saplens hHb3 gene for hair keratin, exons 1 to 9	Homo sapiens mRNA for KIAA1337 protein, partial cds	Homo sapiens mRNA for KIAA1337 protein, partial cds	Homo saplens adaptor-related protein complex AP-4 epsilon subunit mRNA, complete ods	qx57b01.x1 NCI_CGAP_Pan1 Homo sapiens cDNA clone IMAGE:2005417 3'	H.sapiens ECE-1 gene (exon 17)	CM-BT043-090299-075 BT043 Homo saplens cDNA	Homo sapiens DNA for prostacyclin synthase, exon 8	Homo sapiens DNA for prostacyclin synthase, exon 8	Homo sapiens COX11 (yeast) homolog, cytochrome c oxidase assembly protein (COX11), mRNA	Homo sapiens UDP-glucuronosyltransferase 2B4 precursor (UGT2B4) mRNA, UGT2B4*E458 allele,	tridiction and another trial another trial and another trial and another trial and another trial another trial and another trial and another trial and another trial and another trial and another trial and another trial and another trial and another trial and another trial and another trial another trial and another trial and another trial and another trial and another trial and another trial and another trial another trial another trial another trial and another trial another t	Homo sapiens chloride Intracellular channel 4 like (CLIC4L), mkNA	y/74c01.s1 Scares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:248448 3	Homo sapiens T-cell lymphoma invasion and metastasis 1 (TIAM1), mRNA	Homo sapiens Intersectin short isoform (ITSN) mRNA, complete cds	Homo sapiens T-cell lymphoma invasion and metastasis 1 (TIAM1), mRNA	Homo sapiens interesctin short isoform (ITSN) mRNA, complete cds	Human kappa-immunoglobulin germline pseudogene (Chr22.4) variable region (subgroup V kappa II)
Top Hit Acession No.	J94774.1	11417862 NT	/18000.1	4U134963.1	5032192 NT	\B011153.1	4755139 NT	4755139 NT	.76631.1	3F344378.1	3F344378.1	r19208.1	Y19208.1	4B037758.1	4B037758.1	AF155156.2	41263294.1	X91837.1	41904151.1	384122.1	J84122.1	11427788 NT	, 000, 000	41-004200.1	7330334 NT	N59624.1	11526176 NT	AF114488.1	11526176 NT	AF114488.1	M20707.1
Most Similar (Top) Hit BLAST E Value	1.0E-120 ∟	1.0E-120	1.0E-121	1.0E-121	1.0E-121	1.0E-121	1.0E-121	1.0E-121	1.0E-121	1.0E-121	1.0E-121	1.0E-121	1.0E-121	1.0E-121		_		1.0E-121	1.0E-121	1.0E-121	1.0E-121	1.0E-121	707 110 7	1.05-121	1.0E-121	1.0E-121	1.0E-122	1.0E-122	1.0E-122	1.0E-122	1.0E-122
Expression Signal	2.07	1.36	45.	0.98	1.23	0.99	0.89	. 0.89	1.17	1.05	1.05	3.09	3.09	1.09	1.09	19:81	1.21	2.55	1.03	2.58	2.58	4.44		27.78	4.91	3.48	3.99	2.22	1.54	2.85	3.41
ORF SEQ ID NO:		25288		20151		21319	21697	21698										24658	24695	26701	26702					28489	20047	20107	20132		20957
Exan SEQ ID NO:	18722	19111	10052	10328	12674	11461	11818	11818	11945	12404	12404	12989	12969	13408	1	13538	14134	14781	14922	16512	16512	18045				18240	10232	10292	10312	10780	11111
Probe SEQ ID NO:	8914	9495	29	374	707	1556	1923	1923	2055	2530	2530	3042	3042	3492	3492	3624	4236	4901	5050	6632	6632	8167		8163	8336	8363	267	333	355	864	1201

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			_								-	_	_		_	_					_			- 1		4	~	7.00		- FF	——
Top Hit Descriptor	Homo saplens cysteine-rich repeat-containing protein S52 precursor, mRNA, complete cds	Homo sapiens collagen, type XII, alpha 1 (COL12A1), mRNA	Homo sapiens collagen, type XII, alpha 1 (COL;12A1), mRNA	601497032F1 NIH_MGC_70 Homo sapiens cDNA clone IMAGE:3899358 6'	601896173F1 NIH_MGC_19 Homo saplens CDNA clone IMAGE:4125234 5'	601896173F1 NIH_MGC_19 Homo seplens cDNA clone IMAGE:4125234 5	Homo sapiens amylold beta (A4) precursor protein (protease nextn-II, Alzheimer disease) (APP), mRNA	ULHF-BN0-al-a-03-0-UI.1 NIH_MGC_50 Homo sapiens cDNA clone IMAGE:3079948 5'	601113567F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3354232 5	Homo saplens lethal giant larvae (Drosophila) homolog 2 (LLGL2), mRNA	Homo sapiens phosphomannomutase 1 (PMM1), mRNA	Human phosphoendpyruvate carboxykinase (FCK1) gene, promoter region and partial cds	602018058F1 NCI_CGAP_Bm67 Homo sapiens cDNA clone IMAGE:4153670 5	602018058F1 NCI_CGAP_Bm67 Homo sapiens cDNA clone IMAGE:4153670 5'	Homo sapiens chromosome 21 segment HS21C049	Homo sapiens inner membrane protein, mitochondrial (mitofilin) (IMMT), mRNA	Homo sepiens phosphatidylinositol-4-phosphate 5-kinase, type II, beta (PIP5K2B) mRNA, and translated products	Homo saplens phosphatidylinositol-4-phosphate 5-kinase, type II, beta (PIP5K2B) mRNA, and translated	products	Homo sapiens partial mRNA for immunoglobulin kappa chain varlable region (IGVK gene), sample GN02	Human amelogenin (AMELY) gene, 3' end of ¢ds	Human amelogenin (AMELY) gene, 3' end of ¢ds	Human amelogenin (AMELY) gene, 3' end of cds	Homo sapiens RAB9-like protein (LOC51209), mRNA	Homo saplens retinaldehyde-binding protein (CRALBP) gene, complete cds	Homo sapiens retinaldehyde-binding protein (CRALBP) gene, complete cds	601591108F1 NIH_MGC_7 Homo saplens cDNA clone IMAGE:3945433 5'	AU118435 HEMBA1 Homo saplens cDNA clone HEMBA1003591 5'		М	Homo sapiens mRNA for KIAA0454 protein, partial cds
Top Hit Database Source	L	N	TN	EST_HUMAN	EST_HUMAN	EST_HUMAN	Ę	EST_HUMAN	EST_HUMAN	NT	Į.	NT	EST_HUMAN	EST_HUMAN	NT	N	Ę		NT	Ę	LN L	LN L	TN	NT NT	۲N	FN.	EST_HUMAN	EST_HUMAN	N	EST_HUMAN	LN
Top Hit Acession No.	AF167706.1	11418424 NT	11418424 NT	3E906024.1	3F316170.1	3F316170.1	4502166 NT	4W504645.1	BE256039.1	11424216 NT	11418187 NT	J31519.1	BF345274.1	BF345274.1	AL163249.2	5803114 NT	4505818 NT		4505818 NT	AJ388641.1	M55419.1	M55419.1	M55419.1	7705962 NT	L34219.1	L34219.1	BE799746.1	AU118435.1	U42224.1	BE263001.1	AB007923.1
Most Similar (Top) Hit BLAST E Value	1.0E-122	1.0E-122	1.0E-122	1.0E-122	1.0E-122	1.0E-122	1.0E-122	1.0E-122	1.0E-122	1.0E-122	1.0E-122	1.0E-123	1.0E-123	1.0E-123	1.0E-123	1.0E-123	1 OF-123		1.0E-123	1.0E-123	1.0E-123	1.0E-123	1.0E-123	1.0E-123	1.0E-123	1.0E-123	1.0E-123	1.0E-123	1.0E-123		1.0E-123
Expression Signal	1.32	2.01	2.01	5.91	10.56	10.58	1.82	1.2	6.41	1.35	4.35	1.07	1.72	1.72	3.79	5.06	2 34		3.35	1.78	2.98	2.98	2.98	4.6	1.75	1.75	1.29	2.27	1.3	1.95	4.14
ORF SEQ ID NO:	21435	21458				22216	24409		. 25390	27286		19968		L		20767			20977	21211					25091						ŀ
Exon SEQ ID NO:	11669	11586	11588	11871	12318	12318	14623	14797	15337	17096	18869	10153	10881	10681	10916	10923	l	1	11126	11344	1_	11942		_	L	L	L	L	L		<u> </u>
Probe SEQ ID NO:	1667	1684	1684	1772	2441	2441	4738	4918	6051	7219	8608	181	751	751	966	1005	2,0	9171	1218	1439	2052	2052	2052	2267	5344	5344	5425	5893	6186	6059	7392

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Probe	Exan	0		Most Similar		Top Hit	
SEQ ID NO:	SEQ ID NO:	D NO:	Signal	(10p) HIE BLASTE Value	I op hit Acession No.	Dat abase Source	Top Hit Descriptor
7471	17331	27637	12.23	1.0E-123	U09823.1	Ŋ	Oryctolagus cuniculus New Zealand white elonigation factor 1 alpha (Rabefla2) mRNA, complete cds
8928		29057	5	1.0E-123	BF677292.1	EST_HUMAN	602086791F1 NIH_MGC_83 Homo sapiens cDNA clone IMAGE:4250879 5
8968			9	1.0E-123	BF677292.1	EST_HUMAN	602086791F1 NIH_MGC_83 Homo sapiens cDNA clone IMAGE:4250879 5'
268		20048	1.1	1.0E-124	4507500 NT	MT	Homo sapiens T-cell lymphoma invasion and metastasis 1 (TIAM1) mRNA
268			1.1	1.0E-124	1507500	NT	Homo sapiens T-cell lymphoma Invasion and metastasis 1 (TIAM1) mRNA
273	10239		1.98	1.0E-124 D87675.1		뒫	Homo sapians DNA for amyloid precursor protein, complete cds
477	10421	20238	2.11	1.0E-124	1.0E-124 AL163246.2	TN	Homo sapiens chromosome 21 segment HS21 C046
876	10609	20429	2.03	1.0E-124	1.0E-124 AA397551.1	EST_HUMAN	z81b04.r1 Stratagene schizo brain S11 Homolsapiens cDNA clone IMAGE:728719 5' similar to TR:G300482 G300482 POL=REVERSE TRANSCRIPTASE HOMOLOG (RETROVIRAL ELEMENT) ;
676	10609	20430	2.03	1.0E-124	1.0E-124 AA397551.1	EST HUMAN	z81b04.r1 Stratagene schizo brain S11 Homo sapiens cDNA clone IMAGE:728719 6' similar to TR:G300482 G300482 POL=REVERSE TRANSCRIPTASE HOMOLOG (RETROVIRAL ELEMENT) :
742	10673	20509	3.67	1.0E-124		N FA	Human putative ribosomal protein S1 mRNA
791	10720	20561	1.08	1.0E-124	4507500 NT	¥	Homo sapiens T-cell lymphoma invasion and metastasis 1 (TIAM1) mRNA
886	10812		1.81	1.0E-124	7705446 NT	NT.	Homo sapiens hypothetical protein (HSPC088), mRNA
1293			2.59	1.0E-124	11419092	NT	Homo sapiens ring finger protein (RNF), mRNA
1324			6.54	1.0E-124 A		NT	Homo sapiens glucose transporter 3 gene, exons 9, 10, and complete cds
1324		21087	5.54	1.0E-124		NT	Homo sapiens glucose transporter 3 gene, exons 9, 10, and complete cds
1773			2.35	1.0E-124		NT	Homo saplens mRNA for nucleolar RNA-helicase (noH61 gene)
2018			2.23	1.0E-124		EST_HUMAN	601491715F1 NIH_MGC_69 Homo sapiens cDNA clone IMAGE:3893954 5'
2408			0.85	1.0E-124	1.0E-124 AB024069.1	NT	Homo sapiens gene for B120, exon 11
3319			0.85	· 1.0E-124	4504116 NT	ΙN	Homo sapiens glutamate receptor, ionotropic, icalnate 1 (GRIK1) mRNA
3446			96.0	1.0E-124 S78684.1		NT	Homo sapiens ATP-sensitive Inwardly rectifying K-channel subunit (KCNJ6/BIR1) gene, exon
3446			0.86	1.0E-124 S78684.1	78684.1	NT	Homo sapiens ATP-sensitive inwardly rectifying K-channel subunit (KCNJG/BIR1) gene, exon
3598	_		2.95	1.0E-124 X	13794.1	NT	H.sapiens lactate dehydrogenase B gene exon 1 and 2 (EC 1.1.1.27) (and joined CDS)
3826			1.09	1.0E-124	4507500 NT	NT	Homo sapiens T-cell lymphoma invasion and metastasis 1 (TIAM1) mRNA
3983			1.18	1.0E-124	14116	LN.	Homo sapiens glutamate receptor, ionotropic, kainate 1 (GRIK1) mRNA
4639	14527	24315	1.58	1.0E-124	1.0E-124 AB024069.1	NT	Homo sapiens gene for B120, exon 11
4850	14731		1.12	1.0E-124	1.0E-124 M18178.1	TN	Human fibronectin gene extra type III repeat (EDII), exon x+1
5039		24685	2.72	1.0E-124 AI	204535.1	EST_HUMAN	qf86h03.x1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1754069 3'
5240		24934	8.97	1.0E-124	8922337 NT	TN	Homo sapiens hypothetical protein FLJ10300 (FLJ10300), mRNA
5590			6.43	1.0E-124	F696135.1	EST_HUMAN	602124644F1 NIH_MGC_58 Homo sepiens cDNA clone IMAGE:4281635 5'
6185	┖		3.31	1.0E-124 Y	11717.1	NT	M.musculus mRNA for hoxa3 gene
6805	16684	26874	5.66	1.0E-124	4506654 NT	Ł	Homo sapkans ribosomal protein L5 (RPL5) mRNA

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Table 4
Single Exon Probes Expressed in Heart

11/5/2/4																					PCI	/υ	3 0	11/0	JUC)(()
	TR:095162	similar to TR:095162					5,			R:031662 031662	R:031662 031662	nilar to TR:G300482	nilar to TR:G300482						*		3 3' similar to			-	1		0 43
Top Hit Descriptor	hg94a09x1 NCI_CGAP_Kid11 Homo sapien's cDNA cione IMAGE:2953240 3' similar to TR:095162 095162 PEROXISOMAL SHORT-CHAIN AILCOHOL DEHYDROGENASE.	. რ	1	т	П	1	UI-HF-BN0-akz-b-04-0-UI.r1 NIH_MGC_50 Homo sapiens cDNA clone IMAGE:3078846 5	Human muscle glycogen phosphorylase (PYGM) gene, exons 6 through 17	hj05c08.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2980906 3'	#19e03.x1 NCI_CGAP_Gas4 Homo saplens cDNA clone IMAGE:2141980 3' similar to TR:031662 031662 YKRS PROTEIN ;	[f19e03.x1 NCI_CGAP_Gas4 Homo sapiens cDNA clone IMAGE:2141980 3' similar to TR:O31662 O31662 YKRS PROTEIN ;	281b04.r1 Stratagene schizo brain S11 Homo sepiens cDNA clone IMAGE:728719 6' similar to TR:G300482 G300482 POL=REVERSE TRANSCRIPTASE HOMOLOG (RETROVIRAL ELEMENT):	z/81b04.r1 Stratagene schizo brain S11 Homo saplens cDNA clone IMAGE:728719 5' similar to TR:G300482 G300482 POL=REVERSE TRANSCRIPTASE HOMOLOG (RETROVIRAL ELEMENT)	Homo saplens calcineurin binding protein 1 (KIAA0330), mRNA	Homo saplens calcineurin binding protein 1 (KIAA0330), mRNA	Homo saplens mRNA for KIAA1172 protein, partial cds	601577981F1 NIH_MGC_9 Homo saplens cDNA clone IMAGE:3926685 5'	HA0086 Human fetal liver cDNA library Homo sapiens cDNA	HA0086 Human fetal liver cDNA library Homo sapiens cDNA	Homo sapiens ALR-like protein mRNA, partial cds	zk53c07.s1 Soares_pregnant_uterus_NbHPU Homo sapiens cDNA clone IMAGE:486540.3' similar to gb:X85857_cds1 OLFACTORY RECEPTOR*LIKE PROTEIN HGMP07E (HUMAN);	Homo sapiens chromosome 21 segment HS21C010	Homo sapiens KIAA0744 gene product; historie deacetylase 7 (KIAA0744), mRNA	Homo sapiens KIAA0022 gene product (KIAA0022), mRNA	Homo sapiens Usurpin-alpha mRNA, complete cds	Homo sapiens Usurpin-alpha mRNA, complete cds	
Top Hit Database Source	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	Ā	EST HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	EST HUMAN	Ί.	FZ	NT L	EST_HUMAN	EST_HUMAN	EST_HUMAN	IN	EST_HUMAN	NT	NT	NT	LN	Ŋ	
Top Hit Acessian No.	AW612106.1	AW612106.1	345633.1		1.0E-124 AI767133.1	1.0E-124 AI767133.1	AW503755:1	U94776.1	AW665663.1	A1446455.1	1.0E-124 AI446455.1	1.0E-124 AA397651.1	1.0E-124 AA397551.1	11417862 NT	11417862 NT	1.0E-125 AB032998.1	BE743922.1	1.0E-125 AI110656.1	1.0E-125 AI110656.1	AF264750.1	1.0E-125 AA042813.1	1.0E-125 AL163210.2	7662279 NT	7661867	1.0E-125 AF015450.1	1.0E-125 AF016460.1	
Most Similar (Top) Hit BLAST E Value	1.0E-124 AW	1.0E-124 AW	1.0E-124	1.0E-124	1.0E-124	1.0E-124	1.0E-124 AW	1.0E-124 U94776.1	1.0E-124 AW	1.0E-124 AI44	1.0E-124	1.0E-124	1.0E-124	1.0E-124	1.0E-124	1.0E-125	1.0E-125 BE7	1.0E-125	1.0E-125	1.0E-125	1.0E-125	1.0E-125	1.0E-125	1.0E-125	1.0E-125	1.0E-125	
Expression Signal	1.35	1.35	2.44	2.44	7.8	7.8	1.25	2.25	2.25	1.87	1.87	3.88	3.98	1.99	1.99	5.41	3.92	1.63	1.63	1.24	2.13	1.53	1.83	1.08	3.81	3.81	
ORF SEQ ID NO:	27002	27003	27532	27533	27604	27605	27752	28532	28788	28162	28163	20429	20430	25005	25006		19778	20377	20378	20469	20619	20750	20890	21414	21545	21546	
Exon SEQ ID NO:	16808	16808	17326	17326	17393	17393	17526	18280	18509	17916	17916	10609	10609	19628	19626	10278	9987	10565	10585	10643	10769	10905	11050	12700	11668	11668	
Probe SEQ ID NO:	0689	0889	7466	7466	7542	7642	7676	8404	8645	8767	8767	9173	9173	9846	9846	316	420	628	628	711	842	982	1136	1649	1769	1769	

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Single Exon Probes Expressed in Heart

		_																				_				_ +		٠.	• • • • •	r-++ 1
Top Hit Descriptor	2k53c07.s1 Soures, pregnant, uterus, NbHPU Homo sapiens cDNA clone IMAGE:486540 3' similar to gb:X65857, ods.1 OLFACTORY RECEPTOR-LIKE PROTEIN HGMP07E (HUMAN);	Homo sapiens inhibin, alpha (INHA) mRNA	Homo sapiens inhibin, alpha (INHA) mRNA	oh64d02.x5 NCI_CGAP_Kld5 Homo sapiens cDNA clone IMAGE:1471779 3'	Homo sapiens zinc finger protein ZNF287 (ZNF287), mRNA	Homo saplens zinc finger protein ZNF287 (ZNF287), mRNA	601141152F1 NIH_MGC_9 Hamo sapiens cDNA cione IMAGE:3140796 5'	Homo saplens KIAA0985 protein (KIAA0985), mRNA	601433472F1 NIH_MGC_72 Homo sapiens cDNA clone IMAGE:3918952 5	601335826F1 NIH_MGC_44 Homo saplens clDNA clone IMAGE:3689790 5'	601335826F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3689790 5'	Homo sapiens IGF-II gane, exon 5	Homo sapiens IGF-II gene, expn 5	Human chromosome 10 duplicated adrenoleukodystrophy (ALD) gene segment containing exons 8-10	Himan chromosome 10 dinilicated adienoles (2004-chronby (ALD) gene seament containing awas 8-10	QV1-HT0638-070500-191-d12 HT0838 Home sapiens cDNA	QV1-HT0638-070500-191-d12 HT0638 Homo sapiens cDNA	Homo saplens I-REL gene, exon 5	X5900.2X1 NCI_CGAP_Ges4 Homo sepiens cDNA clone IMAGE:2622363 3' similar to TR:Q13284 Q13284 LAMBDA/IOTA PROTEIN KINASE C-INTERACTING PROTEIN [1]:	x85902.x1 NCL_CGAP_Gas4 Homo saplens cDNA clone IMAGE:2622363 3' similar to TR:Q13284 Q13284 LAMBDA/IOTA PROTEIN KINASE C-INTERACTING PROTEIN [1]	Homo saplens mRNA for KIAA0667 protein, partial cds	Homo sapiens myosin, heavy polypeptide 1, skeletal muscle, adult (MYH1), mRNA	Homo saplens poly(A) binding protein II (PABP2) gene, complete cds	RC3-ST0186-250200-018-c11 ST0186 Home sapiens cDNA	QV3-BT0569-020200-075-909 BT0569 Homo sapiens cDNA	QV3-BT0569-020200-075-009 BT0569 Homo saplens cDNA	Homo sapiens mRNA for KIAA0667 protein, partial cds	Homo sapiens CDC-like kinase (CLK) mRNA	Human laminin B1 chain gene, exon 20	
Top Hit Database Source	EST HUMAN	NT	NT	EST_HUMAN	N	NT	EST_HUMAN	N.	EST_HUMAN	EST_HUMAN	EST_HUMAN	F	Z	M	5	EST HUMAN	EST HUMAN	Z	EST_HUMAN	EST HUMAN	1 5	Ę	IN.	EST HUMAN	EST HUMAN	EST_HUMAN	Z	N F	NT	
Top Hit Acession No.	4A042813.1	4504698 NT	4504696 NT	41732966.1	11425114 NT	11425114 NT	3E315412.1	11436448 NT	1.0E-125 BE892660.1	1.0E-125 BE562526.1	1.0E-125 BE562526.1	(03427.1	K03427.1	J80288.1	Ian 288 1	1.0E-125 BE181640.1	1.0E-125 BE181640.1	AF043458.1	4W131202.1	1.0E-125 AW131202.1	4B014567.1	7669505 NT	1.0E-125 AF026029.1	1.0E-125 AW812899.1	1.0E-125 BE074267.1	3E074267.1	4B014567.1	4758007 NT	M61836.1	
Most Similar (Top) Hit BLAST E Value	1.0E-125 AA(1.0E-125	1.0E-125	1.0E-125 AI7	1.0E-125	1.0E-125	1.0E-125 BE	1.0E-125	1.0E-125	1.0E-125	1.0E-125	1.0E-125	1.0E-125 X03427.1	1.0E-125 Ugg	1 0F-125 190288 1	1.0E-125	1.0E-125	1.0E-125 AFC	1.0E-125 AW	1.0E-125	1.0E-125 AB(1.0E-125	1.0E-125	1.0E-125	1.0E-125	1.0E-125 BE	1.0E-126 AB(1.0E-126	1.0E-126 M6	
Expression Signal	1.15	1.65	1.65	2.45	1.98	1.98	0.84	1.41	3.44	1.48	1.48	6.36	6.36	1.22	120	4.31	4.31	3.15	1.86	1.86	6.13	2.92	5.15	2.49	4.36	4.38	1.98	0.88	1.2	
ORF SEQ ID NO:	22228	22316		22320	24136	24137	24195	25573	25610	25993	25994	26093	26094	27043	27044			28211	28289	28290	28616	28748	28754	28804	28896	28897	29044	20525	20528	
Exon SEQ ID NO:	12332	12424	12424	12427	14344	14344	14409	15496	15527	15870	15870	15961	15961	16851	18831	17149	17149	17960	18040	18040	18351	18476	18482	18522	18607	18607	18749	10687	10690	
Probe SEQ ID NO:	2455	2551	2551	2555	4450	4450	4516	5581	5612	5965	5965	6201	6201	6974	6974	7272	7272	8069	8152	8152	8478	8609	8615	8704	8793	8793	8941	757	760	

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Homo sapiens leukocyte immunoglobulin-like receptor, subfamily A (with TM domain), member 1 (LILRA1), чэто sapiens leukocyte immunoglobulin-like receptor, subfamily A (with TM domain), member 1 (LILRA1), 2x86e03.r1 Soares_total_fetus_NbZHF8_9w Homo sapiens cDNA clone IMAGE:796444 5' similar to Human mRNA for cytokeratin 18 zx42a02.r1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA clone IMAGE:789098 5* 2072c03.r1 Stratagene pancreas (#937208) Homo sapiens cDNA clone IMAGE:592420 5/ 2072c03.r1 Stratagene pancreas (#937208) Homo sapiens cDNA clone IMAGE:592420 5/ yx78c06.r1 Soares melanocyte 2NbHM Homo sapiens cDNA clone IMAGE:267850 5' 602139138F1 NIH_MGC_46 Homo saplens cDNA clone IMAGE:4298240 5 601149404F1 NIH_MGC_19 Homo saplens cDNA clone IMAGE:3502129 5 601577981F1 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:3926685 5 Homo sapiens intersectin short isoform (ITSN) mRNA, complete cds Homo sapiens collagen type XI alpha-1 (COL 1/1A1) gene, exon 63 Homo sapiens collagen type XI alpha-1 (COL1/1A1) gene, exon 63 Homo sapiens lost on transformation LOT1 mRNA, complete cds **Top Hit Descriptor** Homo sapiens DNA for amyloid precursor protein, complete cds Homo sapiens DNA for amyloid precursor protein, complete cds Homo sapiens mRNA for casein kinase I epsilon, complete cds Homo sapiens mRNA for casein kinase I epsilion, complete cds Homo sapiens mRNA for casein kinase I epsilon, complete cds Homo sapiens mRNA for casein kinase I epsilon, complete cds Homo sapiens ubiquitin specific protease 8 (USP8) mRNA Homo saplens RAN binding protein 2 (RANBP2), mRNA Homo sapiens mRNA for KIAA1525 protein, partial cds Homo sapiens mRNA for KIAA1525 protein, partial cds Homo saplens ribosomal protein L26 (RPL26) mRNA H.sapiens DNA for liver cytochrome b5 pseudogene H.sapiens gene for alpha1-antichymotrypsin, exon 3 Homo sapiens death receptor 6 (DR6), mRNA Homo sapiens adlican mRNA, complete cds Human mRNA for ankyrin (variant 2.1 TR:G1145880 G1145880 TITIN; **ARNA** mRNA EST_HUMAN NT NT **EST HUMAN** EST HUMAN EST_HUMAN **EST HUMAN** HUMAN EST_HUMAN EST_HUMAN Top Hit Database Source EST 눋 눋 눋 눋 닐눋 5803065 NT 5803065 NT 눋 뉟 4827053 NT 6382078 Top Hit Acession 1.0E-126 AF101108.1 1.0E-126 AF101108.1 1.0E-127 AA450131.1 1.0E-127 AF245505.1 .0E-126 AA460075.1 .0E-128 BE743922.1 .0E-127 AB024597.1 1.0E-126 AA160709.1 1.0E-126 AA160709.1 BE261660.1 I.0E-127 AB024597.1 I.0E-127 AB024597.1 1.0E-127 AF114488.1 .0E-126 AB040958.1 ģ BF683175. 1.0E-126 X53941.1 1.0E-126 7 N34078.1 AB024597 .0E-127 D87675.1 D87675.1 J72621.2 .0E-126 X16809.1 X68735. .0E-126 1.0E-126 1.0E-128 .0E-127 1.0E-127 :0E-127 1.0E-128 1.0E-127 .0E-126 1.0E-126 1.0E-127 (Top) Hit BLAST E **fost Similar** . 1-Value 4.01 4.38 3.59 96.0 7.45 0.96 3.68 3.82 3.59 2.31 2.35 1.32 .28 89 1.59 96'0 2.24 2.04 2.77 2.31 6.07 1.57 1.95 241 8.07 Expression Signal 21939 22075 22330 28910 24890 19954 19954 19955 20640 21800 22341 ORF SEQ ID NO: 20670 22317 22758 24356 24357 25813 25814 26737 19955 20057 21433 22757 23281 23307 24400 10138 12437 SEQ ID 13519 14563 15686 15703 16540 10138 10138 10138 11911 12041 12450 10826 12963 12963 14614 15098 10238 10789 11911 12425 14563 18114 18620 11567 13491 2020 2284 2566 2579 5779 5797 5797 9835 165 166 166 89 89 89 89 2020 Probe SEQ ID NO: 2662 3035 3035 3677 3605 6660 8233 8806 165 4677 4677 4728 272 1665 901 272

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	Top Hit Descriptor	zx42a02.r1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA clone IMAGE:789098 5'	au80e08.y/ Schneider fetal brain 00004 Home sapiens cDNA clone IMAGE:2782594 5' similar to TR:Q16170 Q15170 TRANSCRIPTION FACTOR S-II-RELATED PROTEIN ; contains element MER22	repetitive element;	Homo sapiens delayed rectifier potassium chainnel subunit IsK mRNA, complete cds	Homo sapiens chromosome 21 segment HS21C047	Homo sapiens neuroblastoma-amplified protain (LOC51594), mRNA	Homo sapiens neuroblastoma-amplified protein (LOC51594), mRNA	Homo sapiens cytochrome P450 retinoid metabalizing protein P450RAI-2 mRNA, complete cds	Homo sapiens RAD1 (S. pombe) homolog (RAD1) mRNA, and translated products	Homo sepiens chromosome 21 segment HS21 C068	Homo sapiens Ring1 and YY1 binding protein((RYBP), mRNA	H.saplens NOS2 gene, exon 6	H.saplens TCF11 gene, exon 3-6	Homo sapiens integrin, beta 8 (ITGB8) mRNA.	Homo saplens Pendred syndrome (PDS), mRNA	Homo sapiens Pendred syndrome (PDS), mRNA	Homo sapiens secretory pathway component Sec31B-1 mRNA, alternatively spliced, complete cds	Homo saplens secretory pathway component Sec31B-1 mRNA, alternatively spliced, complete cds	Homo sapiens Chediak-Higashi syndrome 1 (CHS1), mRNA	Homo sapiens similar to heat shock 70kD protein 9B (mortalin-2) (H. sapiens) (LOC63184), mRNA	Homo sapiens similar to heat shock 70kD protein 9B (mortalin-2) (H. sapiens) (LOC63184), mRNA	601434784F1 NIH_MGC_72 Homo saplens cDNA clone IMAGE:3918917 5	601434784F1 NIH_MGC_72 Homo sapiens cDNA clone IMAGE:3919917 5	Homo sapiens mRNA for casein kinase I epsilon, complete ods	Hamo sapiens mRNA for casein kinase I epsilon, complete cds	Hamo eapiens gene for AF-6, complete cds	601278127F1 NIH_MGC_20 Homo sapiens cDNA clone IMAGE:3618822 5'	Homo saplens chondroitin sulfate proteoglycen 2 (versican) (CSPG2) mRNA	Homo sapiens chondroitin sulfate proteoglycen 2 (versican) (CSPG2) mRNA	Human FAU1P pseudogene, trinuclectide repeat regions	Human FAU1P pseudogene, trinucleotide repeat regions
	Top Hit Database Source	EST HUMAN		EST_HUMAN	NT	NT	NT	NT	NT	NT	NT	Ľ	N	Ł	LN-	NT	NT	TN	TN	LN	NT	NT	EST_HUMAN	EST_HUMAN	LN	NT	NT	EST_HUMAN	NT	LN	NT	NT
	Top Hit Acessian No.			1.0E-127 AW161297.1		163247.2	7706239 NT	TN6239 NT	F262297.1	4506384 NT	163268.2	6912639 NT			4504778 NT	11421914 NT	11421914 NT	1.0E-127 AF274863.1	AF274863.1	11427235 NT	11417339 NT	11417339 NT	3E895415.1	1.0E-127 BE895415.1	1.0E-127 AB024597.1	1.0E-127 AB024597.1	1.0E-127 AB011399.1	1.0E-128 BE385617.1	4758081 NT	4758081 NT	J02523.1	J02523.1
	Most Similar (Top) Hit BLAST E Value	1.0E-127 AA450131.1		1.0E-127	1.0E-127 AI	1.0E-127 AI	1.0E-127	1.0E-127	1.0E-127 A	1.0E-127	1.0E-127 AI	1.0E-127	1.0E-127 X85764.1	1.0E-127 X84060.1	1.0E-127	1.0E-127	1.0E-127	1.0E-127	1.0E-127	1.0E-127	1.0E-127	1.0E-127	1.0E-127 B	1.0E-127			1.0E-127	1.0E-128	1.0E-128	1.0E-128	1.0E-128	1.0E-128 U02523.1
	Expression Signal	0.98		0.88	1.09	0.86	21.46	21.48	0.92	4.16	1.93	96.0	3.72	2.67	6.76	1.38	1.38	4.97	4 97	1.17	6.74	6.74	2.46	2.48	1.68	1.66	2.1	4.46	1.48	1.48	12.19	12.19
	ORF SEQ ID NO:	22342		23435			23833	L			L	24252						27624				L				19955	25276			20893		21806
	Exan SEQ ID NO:	12450		13652	L		L		1		1	1		1		L		1		┸				18707	L	乚	19177	١.	1	ı	ı	11916
	Probe SEQ ID NO:	2579		3740	4028	4128	4160	4160	4395	4489	4532	4575	6548	5742	5816	6578	6578	7558	7558	7923	8498	8498	8898	8888	2397	8397	9595	452	1138	1138	2025	2025

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Table 4
Single Exon Probes Expressed in Heart,

																	-1			"r				n29		n.	JI TL.	2.1178
Top Hit Descriptor	Homo sapiens ribosomal protein S2 (RPS2) mRNA	Homo saplens chromatin-specific transcription elongation factor, 140 kDa subunit (FACTP140), mRNA	Homo sapiens mRNA for KIAA1247 protein, partial cds	Homo sapiens prospero-related homeobox 1 (PROX1), mRNA	Homo saplens phosphodlesterase 1C, calmodulin-dependent (70kD) (PDE1C), mRNA	7q86b10.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE: 3'	Homo sapiens mRNA for KIAA0454 protein, partial cds	Homo sapiens mRNA for KIAA0454 protein, pertial cds	ns04a11.11 NOI_CGAP_Ew1 Homo saplens cDNA clone IMAGE:1182820 similar to TR:G951338 G951338 CHROMOSOME SEGREGATION GENE HOMOLOG CAS.	Homo sapiens glutamate receptor, lonotropic, N-methyl D-aspartate 2D (GRIN2D), mRNA	om88h08.s1 NCI_CGAP_GC4 Homo saplens cDNA clone IMAGE:1552383 3' similar to gb:X54941 CYCLIN- DEPENDENT KINASES REGULATORY SUBUNIT 1 (HUMAN);	801277828F1 NIH MGC 20 Homo saplens cDNA clone IMAGE:3818750 5'	EST367360 MAGE resequences, MAGC Homo sapiens cDNA	insulin-like growth factor binding protein-2 [human, placenta, Genomic, 1019 nt, segment 2 of 4]	Insulin-like growth factor binding protein-2 [human, placenta, Genomic, 1019 nt, segment 2 of 4]	Novel human mRNA containing Zinc finger C2H2 type domains	Homo seplens glutathione S-transferase theta 2 (GSTT2) and glutathione S-transferase theta 1 (GSTT1) genes, complete cds	Homo saplens glutathione S-transferase theta 2 (GSTT2) and glutathione S-transferase theta 1 (GSTT1) dense. complete cds	Homo saplens zinc finger protein 76 (expressed in testis) (ZNF76), mRNA	Homo sapiens platelet-derived growth factor receptor, beta polypeptide (PDGFRB) mRNA	Homo sapiens platelet-derived growth factor receptor, beta polypeptide (PDGFRB) mRNA	ZINC FINGER PROTEIN HZF10	ZINC FINGER PROTEIN HZF10	ZINC FINGER PROTEIN HZF10	Homo sapiens mRNA for KIAA1459 protein, partial cds	CMYA5 Human cardiac muscle expression library Homo saplens cDNA clone 4151935 similar to CMYA5	Cardiomyopathy associated gene 5	CMYA5 Human cardiac muscle expression libiary Homo sapiens cDNA clone 4151935 similar to CMYA5 Cardiomyopathy associated gene 5
Top Hit Database Source	FN	Į	ZZ FZ	N	ĘZ	EST_HUMAN	NT	NT	EST_HUMAN	ΙΝ	EST HUMAN	EST HUMAN	EST HUMAN	LN	NT	TN	Į,	F	LZ	FZ	닐	SWISSPROT	SWISSPROT	SWISSPROT	NT		EST_HUMAN	EST_HUMAN
Top Hit Acession No.	4506718 NT	11437455 NT	8	11426673 NT	11420965 NT	1.0E-128 BF224345.1			1.0E-128 AA639198.1	11425254 NT	1.0E-128 AA926959.1	3E384475.1	4W955280.1	537722.1	537722.1	1.0E-129 AL096880.1	1.0E-129 AF240786.1		18522	4505682 NT	4505682 NT				32.1		4W 755254.1	755254.1
Most Similar (Top) Hit BLAST E Vatue	1.0E-128	1.0E-128	1.0E-128 AB0	1.0E-128	1.0E-128	1.0E-128	1.0E-128	1.0E-128	1.0E-128	1.0E-128	1.0E-128	1.0E-128 BE3	1.0E-128 AW	ĺ			1.0E-129	1 0F-129	1.0E-129	1.0E-129	1.0E-129	1.0E-129 Q14585	1.0E-129	1.0E-129 Q14585	1.0E-129		1.0E-129 AW	1.0E-129 AW
Expression Signal	13.3	0.85	1.13	5.48	2.67	7.23	3.28	3.28	1.25	5.94	3.87	1.79	3.66	2.18	1.35	2.86	2.29	2.29	2.43	1.19	1.19	1.35	1.35	1.35	2.2		9.7	9.7
ORF SEQ ID NO:	21850		23071	24245	25830	26123	27046	27047	27929	28232	28240	28364		20182	20182	21461	21465	21468	21575	22503	22504	22808	22809	22810	23764		23856	23857
Exon SEQ ID NO:	12049	12273	13268	<u>L</u> _	15771	15988		16853	17684	17983	17991	18111	18968	10353	10353	11591	11595	11595	L	12613	12613				13975		14083	14083
Probe SEQ ID NO:	2162	2395	3348	4565	5865	6140	6976	6976	7834	8092	8101	8230	9263	116	407	1689	1693	1693	1802	2751	2751	3089	3089	3089	4073		4183	4183

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Single Exon Probes Expressed in Heart

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
2706	15814	25715	289	1.0E-129	AJ006345.1	뉟	Homo sapiens KVLQT1 gene
6229	16095	26245	5.17	1.0E-129	AJ008345.1	뒫	Homo saplens KVLQT1 gene
6267	16132	26286	7.59	1.0E-129	11420850 NT	ΙΝ	Homo sapiens similar to ribosomal protein S26 (H. sapiens) (LOC63694), mRNA
6841	16720		3.68	1.0E-129	AB0145	F	Homo sapiens mRNA for KIAA0634 protein, partial cds
8554	18424	28693	3.52	1.0E-129	AA6255	П	af72f07.r1 Soares_NhHMPu_S1 Homo saplens cDNA clone IMAGE:1047589 5'
8618	16132	26286	9.4	1.0E-129	11420850	11420850 NT	Homo sapiens similar to ribosomal protein S26 (H. sapiens) (LOC63694), mRNA
8888	18699	28993	2.04	1.0E-129	AU143115.1	Г	AU143115 Y79AA1 Homo sapiens cDNA clone Y79AA1001410 5'
8888	18699	28994	2.04	1.0E-129	AU143115.1	EST_HUMAN	AU143115 Y79AA1 Homo saplens cDNA clorie Y79AA1001410 5'
9250	18960		1.87	4 OE-130	1182455 4	ECT LIMAN	yqd9c05.r1 Soares fetal liver spleen 1NFLS Home sapiens cDNA clone IMAGE:199112 6' similar to SDBaaren Baaren Ho 25-Higebnation bei aten decited i tamine acidatic is-acida
9630	19203		1.88	1.0E-129	AL 120739.1	Т	DKFZ0762K171 r1 782 (synonym: hmel2) Hemo sagiens CDNA clone DKFZ0762K171 5'
1643	11547	21408	684	1 0E-130	RE275192 1	Π	601121995F1 NIH MGC 20 Homo saciens cDNA clone IMAGE:3348368 5
1643	11547	21409	6.81	1.0E-130			601121996F1 NIH MGC 20 Homo septens cDNA clane IMAGE:3346366 5
1940	11835		2.06	1.0E-130			Human gene for catalase (EC 1.11.1.6) exon 9 mapping to chromosome 11, band p13
2743	12605		7.7	1.0E-130	AJ010230.1	Ę	Homo saplens RET finger protein-like 1 antisense transcript, partial
2849	12777	22564	1.1	1.0E-130	BE564219.1	EST_HUMAN	601343016F1 NIH_MGC_53 Homo sapiens cDNA clone IMAGE:3685466 5'
2849	12777	22565	1.1	1.0E-130	BE564219.1	EST_HUMAN	601343016F1 NIH_MGC_53 Homo sapiens cDNA clone IMAGE:3685466 5'
3530	13446	23243	1.07	1.0E-130			Homo sapiens retind dehydrogenase homolog isoform-1 (RDH) mRNA, complete cds
3703	12777	22564	4.77	1.0E-130		HUMAN	601343016F1 NIH_MGC_53 Homo saplens cDNA clone IMAGE:3685466 5'
3703	12777	22565	4.77	1.0E-130	BE564219.1	HUMAN	601343016F1 NIH_MGC_53 Homo sapiens cDNA clone IMAGE:3885468 5'
3857	13768	23560	1.09	1.0E-130	AW 503580.1		UI-HF-BN0-aky-g-06-0-UI.r1 NIH_MGC_50 Homo saplens cDNA clone IMAGE:3078731 5'
4438	14333	24122	7.48	1.0E-130	AW843993.1		CM4-CN0045-180200-511-f02 CN0045 Homp sapiens cDNA
6028	14902	24673	1.09	1.0E-130	AW363299.1	EST_HUMAN	RC0-CT0318-201199-031-a11 CT0318 Homo saplens cDNA
5029	14902	24874	1.09	1.0E-130	AW363299.1	EST_HUMAN	RC0-CT0318-201199-031-a11 CT0318 Homo sapiens cDNA
6301	16165	26322	2.04	1.0E-130	11416777 NT	Ę	Homo sapiens solute carrier family 6 (neurotransmitter transporter, L-proline), member 7 (SLC6A7), mRNA
7119	16998	27187	2.45	1.0E-130	AW956242.1	EST_HUMAN	EST368312 MAGE resequences, MAGD Homo saplens cDNA
7314	17180	27392	1.57	1.0E-130	AB037756.1	ΙN	Homo sapiens mRNA for KIAA1335 protein, partial cds
8513	18385	28650	32.43	1.0E-130	M25140.1	NT	Human cardiac alpha-myosin heavy chain (MYH6) gene, exons 2, 3 and 4
4	9991	19782	2.49	0.0E+00	AA228126.1	EST_HUMAN	zr68c04.r1 Soares_NhHMPu_S1 Homo saplens cDNA clone IMAGE:667590 5' similar to TR:G222811 G222811 ALPHA 1 CHAIN OF TYPE XII COLLAGEN ;
.4	9991	19783	2.49	0.0E+00	AA22812		z 58c04.r1 Soares_NhHMPu_S1 Homo sapléns cDNA clone IMAGE:097590 5' similar to TR:G222811 G222811 ALPHA 1 CHAIN OF TYPE XII COLLAGEN;
7	9993	19786	1.44	0.0E+00	4885136	4885136 NT	Homo sapiens checkpoint suppressor 1 (CHES1), mRNA

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	_	_			_	_	_	_	_	_	_		_	_		_		_	,	,	4	+	- 79		(111)	_	1	1:25	•	11.11	1. 17-11-2
Top Hit Descriptor	Homo sapians hypothetical protein FLJ20371 (FLJ20371), mRNA	Homo sapiens hypothetical protein FLJ20371 (FLJ20371), mRNA	Homo saplens DCRR1 mRNA, partial cds	Homo saplens DCRR1 mRNA, partial cds	Homo sapiens beta-tubulin mRNA, complete cds	Human heparin cofactor II (HCF2) gene, exons 1 through 5	Homo saplens RNA-binding protein S1, seriné-rich domain (RNPS1), mRNA	Homo sapiens mRNA for multidrug resistance protein 3 (ABCC3)	Homo sapiens mRNA for multidrug resistance protein 3 (ABCC3)	HUM516H08B Human placenta polyA+ (TFuljwara) Homo sapiens cDNA clone GEN-516H08 5'	HUM516H08B Human placenta polyA+ (TFuliwara) Homo sapiens cDNA clone GEN-516H08 5'	Human ribosomal protein L7 (RPL7) mRNA, complete cds	cr48e07.x1 Jla bane marrow strama Hamo sapiens cDNA clane HBMSC_ar48e07 3'	cr48e07.x1 Jia bone marrow stroma Homo saptens cDNA clone HBMSC_cr48e07.3'	Human von Willebrand factor pseudogene corresponding to exons 23 through 34	Human von Willebrand factor pseudogene corresponding to exons 23 through 34	Homo sapiens protein tyrosine phosphatase, non-receptor type substrate 1 (PTPNS1) mRNA	Homo sapiens protein tyrosine phosphatase, non-receptor type substrate 1 (PTPNS1) mRNA	Homo sapiens protein tyrosine phosphatase, non-receptor type substrate 1 (PTPNS1) mRNA	Homo saplens protein tyrosine phosphatase, rion-receptor type substrate 1 (PTPNS1) mRNA	on89e04.s1 Soares_NFL_T_GBC_S1 Homo septens cDNA clone IMAGE:1563870 3' similar to SW:TMOD_HUMAN P28289 TROPOMODULIN ;	Homo sapiens amiloride binding protein 1 (amine oxidase (copper-containing)) (ABP1), nuclear gene	encoding mitochondrial protein, mRNA	Homo saplens heterogeneous nuclear ribonucleoprotein A1 (HNRPA1) mRNA	Homo sapiens actin, beta (ACTB) mRNA	Human polyhomeotic 1 homolog (HPH1) mRNA, partial cds	HA1347 Human fetal liver cDNA library Homo saplens cDNA	Homo saplens mRNA for KIAA1363 protein, partial cds	H.saptens nox1 gene (exon 2)	ts38b05.x1 NCI_CGAP_Ut4 Homo sapiens cJNA clone IMAGE:2230833 3' similar to TR:Q99551 Q99551 MITOCHONDRIAL TRANSCRIPTION TERMINATION FACTOR PRECURSOR.;	ts38b05.x1 NC_CGAP_Ut4 Homo sapiens cJNA clone IMAGE:2230833 3' similar to TR:Q99551 Q99551 MITOCHONDRIAL TRANSCRIPTION TERMINATION FACTOR PRECURSOR.;
Top Hit Database Source	TN	NT	N T	NT	Z,	ΤΝ	NT	LN LN	INT	EST_HUMAN	EST_HUMAN	NT	EST_HUMAN	EST_HUMAN	Z	Ę	TN	TN	L	FN	EST_HUMAN		NT	NT	NT	NT	EST_HUMAN	NT	ΙN	EST HUMAN	EST_HUMAN
Top Hit Acession No.	8923349 NT	8923349 NT	383327.1	383327.1	4F141349.1	M58600.1	6857825 NT	117151.2	Y17151.2	J78804.1	J78804.1	_16558.1	4W069534.1	4W069534.1	M60676.1	V60676.1	4758977 NT	4758977 NT	4758977 NT	4758977 NT	4A953770.1		4501850 NT	450444 NT	5016088 NT	J89277.1	4/114743.1	AB037784.1	X91213.1	AI623701.1	0.0E+00 AI623701.1
Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	00+30.0 .	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00
Expression Signal	1.34	1.34	2.45	2.45	5.57	26.0	2.41	1.4	1.4	1.22	1.22	4.14	8.1	8.1	5.8	2.75	1.77	1.77	1.49	1.49	42.13		1.09	14.77	47.55	13.39	1.46	1.03	6.13	1.30	1.88
ORF SEQ ID NO:	19792	19793	19789	19800	19804	19816		19845		19847	19848			19853	19857		19869	19870	19869	19870	19875		19877		19886	19889	19895	19896	19901	19906	19906
Exon SEQ ID NO:	10001	10001		10001	10011		10022	10038		10039	10039			10042	10045	10047	10054	10054	10054	10054	10058			10061	10070	10073	10079	10080	10085	10091	10091
Probe SEQ ID NO:	15	15	20	20	24	33	36	51	51	62	52	63	99	65	69	61	69	69	71	71	74		76	77	86	88	94	36	102	110	111

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		_	_					_	_	_			_				_	_	_		100		٠.,				_	 	100		n n.o.	ų
	Top Hit Descriptor	yy01h09.r1 Soares melanocyte 2NbHM Homo(sapiens cDNA clone IMAGE:270017 5'	yy01h09.r1 Soares melanocyte 2NbHM Homo sapiens cDNA clone IMAGE:270017 5'	Homo sapiens neuropilin 2 (NRP2) mRNA	Homo sapiens polymerase (RNA) II (DNA directed) polypeptide A (220kD) (POLR2A) mRNA	Homo sapiens polymerase (RNA) II (DNA directed) polypeptide A (220kD) (POLR2A) mRNA	ya83g04.r2 Stratagene fetal spleen (#337205) Homo saplens cDNA clone IMAGE:68310 5'	ya83g04.r2 Stratagene fetal spleen (#937205) Homo sapiens cDNA clone IMAGE:68310 5'	Homo sapiens heterogeneous nuclear ribonucleoprotein A1 (HNRPA1) mRNA	601460375F1 NIH_MGC_66 Homo sapiens cDNA done IMAGE:3863803 5'	Homo sapiens heterogeneous nuclear ribonucleoprotein A1 (HNRPA1) mRNA	Homo sapiens serine palmitoy/ transferase, subunit II gene, complete cds; and unknown genes	601174270F1 NIH_MGC_17 Homo sapiens cDNA done IMAGE:3529864 5'	601174270F1 NIH_MGC_17 Homo sapiens cDNA done IMAGE:3529864 5'	zd62b05.r1 Soares_fetal_heart_NbHH19W Homo sapiens cDNA clone IMAGE:345201 5' sImilar to	gb:X16282_cds1 ZINC FINGER PROTEIN CLONE 647 (HUMAN);	Homo sapiens zinc finger protein mRNA, complete cds	Homo sapiens chromosome 21 segment HS2i C002	Homo sapiens chromosome 21 segment HS2! C002	bb24e12.y1 NIH_MGC_14 Homo saplens cDNA clone IMAGE:2863854 5' similar to WP:Y57A10A.Z CE22631;	bb24e12.y1 NIH_MGC_14 Homo sepiens cDNA clone IMAGE:2963854 5' similar to WP:Y57A10A.Z	CE22631;	Homo sapiens mRNA for KIAA0784 protein, partial cds	Homo sapiens mRNA for KIAA0784 protein, partial cds	Human gamma-cytoplasmic actin (ACTGP9) pseudogene	Homo sapiens CTCL tumor antigen se14-3 mRNA, complete cds	Homo sapiens CTCL tumor antigen se14-3 mRNA, complete cds	Homo sapiens chromosome XMSL3-2 protein mRNA, complete cds	Homo sapiens chromosome X MSL3-2 protein mRNA, complete cds	tq04f08x1 NCI_CGAP_Ut3 Homo sapiens cDNA clone IMAGE:2207847 3' similar to gb:J03191 PROFILIN I (HUMAN);	tq04f08.x1 NCI_CGAP_US Homo sapiens cDNA done IMAGE:2207847 3' similar to gb:J03191 PROFILIN I (HUMAN):	
	Top Hit Database Source	EST_HUMAN	EST_HUMAN	·			T_HUMAN	EST HUMAN		T_HUMAN			EST_HUMAN	Г		T_HUMAN			F	EST HUMAN	Г	EST_HUMAN	_ LN	I	IN	LN LN	FN	Į.	Į.	EST HUMAN		T
	Top Hit Acession No.	N36040.1	\36040.1	4505458 NT	4505938 NT	4505938 NT	T56945.1	T56945.1	450444 NT	0.0E+00 BF036881.1	4504444 NT	0.0E+00 AF111168.2	0.0E+00 BE295973.1				.1	0.0E+00 AL163202.2	0.0E+00 AL163202.2	0.0E+00 BE018970.1		0.0E+00 BE018970.1	0.0E+00 AB018327.1	0.0E+00 AB018327.1		0.0E+00 AF273045.1	0.0E+00 AF273045.1	Г	Γ	41587308.1	587308.1	
Most Similar	(Top) Hit BLAST E Value	0.0E+00 N36040.1	0.0E+00 N36040.1	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00 W 73973.1	0.0E+00	0.0E+00	0.0E+00/	0.0E+00		0.0E+00	0.0E+00	0.05+00	0.0E+00 D50859.1	0.0E+00	0.0E+00	0.0E+00	0.0E+00/	0.0E+00 A	0.0E+00 A	22.5
	Expression Signal	1.83	1.83	0.86	3.17	3.17	1.49	1.49	9.05	242	15.84	1.36	1.1	1.18		2.99	1.51	18.37	18.37	4.25		4.25	1.98	1.98	130.42	2.83	2.83	2:92	2.92	8.33	9.33	
	ORF SEQ ID NO:	19907	19908	18913	19921	19922	18929	19930	-	18943	<u> </u>	19948	18947	19947		19948	19949	19952	19953	19960		19961	19964	19965	19976	19981	19982	19984	19985	19991	19892	
	SEQ ID NO:	12636	12636	10094	10100		10108	10108	10121	10125	10127	10130	10132		L	10133	10134	10137	10137	10145		10145	10150	10160	10159	10164	10164	10166	<u>l</u> _		ŀ	ł
	Probe SEQ ID NO:	112	112	115	128	126	135	135	147	151	153	166	158	159		160	181	164	164	174		174	179	179	187	192	192	194	194	203	203	

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rCBAP1E4466 Pediatric pre-B cell acute lymphoblastic laukemia Baylor-HGSC project=TCBA Homo saplens CBAP1E4466 Pediatric pre-B cell acute lymphoblastic leukemia Baylor-HGSC project=TCBA Homo saplens CBAP1E4488 Pediatric pre-B cell acute lymphoblastic leukemia Baylor-HGSC project=TCBA Homo sapiens CBAP1E4468 Pediatric pre-B cell acute lymphoblastic leukemia Baylor-HGSC project=TCBA Homo saplens CBAP1E4466 Pediatric pre-B cell acute lymphoblastic leukemia Baylor-HGSC project=TCBA Homo sapiens CBAP1E4466 Pediatric pre-B cell acute lymphoblastic leukemla Baylor-HGSC project=TCBA Homo sapiens CBAP1E4466 Pediatric pre-B cell acute lymphoblastic leukemia Baylor-HGSC project=TCBA Homo sapien: CBAP1E4466 Pediatric pre-B cell acute lymphoblastic leukemia Baylor-HGSC project=TCBA Homo sapien: CBAP1E4466 Pediatric pre-B cell acute lymphoblastic leukemia Baylor-HGSC project=TCBA Homo sapien Homo sapiens T-cell lymphoma invasion and metastasis 1 (TIAM1) mRNA Homo sepiens T-cell lymphoma invasion and metastasis 1 (TIAM1) mRNA Mus musculus testis-specific protein, Y-encoded-like (Tspyl), mRNA **Fop Hit Descriptor** Homo sapiens chromosome 21 unknown mRNA H.sapiens mRNA for Interferon alphe/beta receptor (long form) Homo saptens DCRR1 mRNA, partial cds Homo saptiens DCRR1 mRNA, partial cds IL2-CT0031-181199-020-B03 CT0031 Homo sapiens cDNA Homo sapiens NS1-associated protein 1 (NSAP1) mRNA Homo sapiens hypothetical protein (LOC51250), mRNA Homo sapiens mRNA for KIAA0721 protein, partial cds Homo sapiens mRNA for KIAA0721 protein, partial cds protein L31 (RPL31) mRNA Homo saplens TADA1 protein mRNA, complete cds Homo sapiens chromosome 21 unknown mRNA Homo sapiens chromosome 21 segment HS21 cDNA clone TCBAP4466 cDNA clone TCBAP4466 cDNA clone TCBAP4466 cDNA clone TCBAP4466 cDNA clone TCBAP4468 cDNA clone TCBAP4466 cDNA clone TCBAP4466 cDNA clone TCBAP4468 cDNA clone TCBAP4466 Homo sapiens ribosomal EST_HUMAN EST HUMAN EST_HUMAN EST_HUMAN **EST HUMAN** HUMAN EST_HUMAN **EST_HUMAN** EST_HUMAN EST_HUMAN Top Hit Database Source EST_ Ę 눋 뉟 눋 뉟 눋 눋 눋 4507500 6678444 4506632 Top Hit Acession 0.0E+00 AL163201.2 0.0E+00 AF231919.1 0.0E+00 X89772.1 4507 0.0E+00 AW845293.1 0.0E+00 AF231919.1 0.0E+00 BE246780.1 AF132000.1 0.0E+00 AB018284.1 0.0E+00 66 0.0E+00 BE246780.1 0.0E+00 BE246780.1 0.0E+00 BE246780.1 0.0E+00 BE246780.1 0.0E+00 BE246780.1 BE246780.1 0.0E+00 BE246780.1 0.0E+00 BE246780.1 AB018264.1 ģ 0.0E+00 D83327.1 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 (ΤΦ) Hit BLAST E dost Similar Value 6.54 3.99 3.99 3.99 12.62 12.62 4.66 1.46 6.8 1.14 1.14 2.57 0.86 16.66 2.48 3.43 3.43 3.46 1.95 3.43 Expression Signal 20050 20051 20053 20066 20087 ORF SEQ ID NO: 19999 19999 20002 20005 20006 20004 20005 20032 20004 20006 20004 10186 10195 10213 10215 10234 10234 10236 10246 10246 10179 10186 10195 10195 10195 10195 10195 10195 10195 10180 10195 10223 SEO ID 10187 249 269 269 270 **88** 88 Probe SEQ ID 208 209 215 281 216 226 226 224 224 224 225 225 225 ö

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Single Exon Probes Expressed in Heart

Homo sapiens myeloid/ymphoid or mixed-lineage leukemla (trithorax (Drosophila) homolog); translocated to, Homo sapiens phosphoribosylglycinamide formyltransferase, phosphoribosylglycinamide synthetase, qy81h05.x1 NCL_CGAP_Brn25 Homo sapiens cDNA clone IMAGE:2018457 3' similar to gb:X54199 Homo sapiens potassium inwardly-rectifying channel, subfamily J, member 16 (KCNJ15) mRNA TRANSCRIPTION REGULATOR PROTEIN[BACH1 (BTB AND CNC HOMOLOG 1) (HA2303) TRANSCRIPTION REGULATOR PROTEIN[BACH1 (BTB AND CNC HOMOLOG 1) (HA2303) Homo sapiens polassium inwardly-rectifying channel, subfamily J, member 15 (KCNJ15) mRNA Homo sepiens GA-binding protein transcription factor, alpha subunit (60kD) (GABPA), mRNA Human mRNA for KIAA0184 gene, partial cds Homo saplens hormonally upregulated neu tumor-associated kinase (HUNK), mRNA Homo sapiens hormonally upregulated neu tumor-associated kinase (HUNK), mRNA 2v18c08 r1 Soares_NhHMiPu_S1 Homo sapiens cDNA clone IMAGE:753994 6'
Homo sapiens SON DNA binding protein (SON) mRNA
Homo sapiens SON DNA binding protein (SON) mRNA Homo sapiens T-cell lymphoma invasion and metastasis 1 (TIAM1) mRNA Homo sapiens intersectin short isoform (ITSN) mRNA, complete cds Human mRNA for KIAA0184 gene, partial cds AU134963 PLACE1 Homo sapiens cDNA done PLACE1000899 5 Fop Hit Descriptor RC2-CT0320-300100-016-a09 CT0320 Homo sepiens cDNA Homo seplens IgG Fc binding protein (FC(GAMMA)BP) mRNA Homo sapiens IgG Fc binding protein (FC(GA,MMA)BP) mRNA PHOSPHORIBOSYLAMINE-GLYCINE LIGASE (HUMAN); phosphoribosylaminolmidazole synthetase (GART) mRNA Human zinc finger protein zfp31 (zf31) mRNA, partial cds Homo sapiens mRNA for KIAA1019 protein, partial cds Homo sapiens mRNA for KIAA1019 protein, partial cds Homo sapiens mRNA for KIAA1019 protein, partial cds FRANSCRIPTION REGULATOR PROTEIN BACH1 Homo sapiens X-box binding protein 1 (XBP1) mRNA Homo sapiens chromosome 21 unknown mRNA Homo sapiens chromosome 21 unknown mRNA Homo sapiens chromosome 21 unknown mRNA Homo sapiens moesin (MSN), mRNA (MLLT4) mRNA HUMAN Top Hit Database Source SWISSPROT EST_HUMAN EST_HUMAN SWISSPROT 4557029 NT 4567029 NT 눋 4503914 NT 4503680 NT 4503680 NT Z 눋 4507152 NT 4507152 NT 눌 7857213 NT 7657213 NT 5174574 NT 4827057 NT Ξ E Ł 4506728 4505256 Top Hit Acession 4507500 4503854 0.0E+00 AB028942.1 0.0E+00 AB028942.1 0.0E+00 AW754180.1 0.0E+00 AU134963.1 0.0E+00 AB028942.1 AF231919.1 AF114488.1 0.0E+00 AF231919.1 0.0E+00 AI363014.1 ģ 0.0E+00 AF231919.1 0.0E+00 45 D80008.1 **U71600.1** 0.0E+00 D80006. 0.0E+00 O14867 0.0E+00 L 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.05+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 Aost Similar (Top) Hit BLAST E Value .4.03 13.28 5.26 5.26 9.68 3.11 5.3 96: 0.9 4.18 2.31 0.86 8.0 2.15 2.15 0.89 3.67 3.4 1.52 3.43 1.38 9. Expression Signal 20074 20085 20102 20103 ORF SEQ 20087 20088 20092 20103 20115 20129 20130 20133 20188 20189 20159 20116 20120 20125 20160 20131 20136 20137 20137 Ö Ö Ö 10254 10265 10266 SEQ ID 12664 10267 10269 10386 10336 10268 10285 10285 10286 10286 10300 10313 10315 10316 10318 10273 12665 10326 183<u>4</u> 10307 10311 10365 Š 8 Probe SEQ ID 280 ន្ត 301 302 303 8 305 307 325 325 325 326 341 342 345 353 372 387 387 354 356 369 358 ë

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Single Exon Probes Expressed in Heart

Expn NO: 10339 ORF SEQ 2010 2016 Expression Signal (Top) Hit LAST E NO: 10339 Top Hit Acession Value Top Hit Acession NO: 10339 Top Hit Acession NO: 10339 Top Hit Acession NO: 10339 Top Hit Acession NO: 10339 Top Hit Acession NO: 10339 Top Hit Acession Across NO: 10339 Top Hit Acession Across NO: 10339 Top Hit Acession Across NO: 10340 <th></th> <th></th> <th>_</th> <th></th> <th>_</th> <th>, .</th> <th></th> <th></th> <th></th> <th></th> <th></th> <th>,</th> <th></th> <th></th> <th>,</th> <th>_</th> <th></th> <th>_</th> <th>_</th> <th></th> <th>n</th> <th>٠,,,</th> <th>!!</th> <th></th> <th></th> <th></th> <th></th> <th></th> <th>1.1</th> <th>بيلتم</th> <th><u></u></th> <th>1</th> <th>111.</th> <th>11.70</th> <th>ים ונכ</th>			_		_	, .						,			,	_		_	_		n	٠,,,	!!						1.1	بيلتم	<u></u>	1	111.	11.70	ים ונכ
Even NO: 10339 CRF SEQ D NO: 10339 Expression Signal D NO: 10339 Top Hit Accession Value NO: 10339 Top Hit Accession Accossion NO: 10340 Top Hit Accession Accossion NO: 10340 Top Hit Accession Accossion NO: 10340 Top Hit Accession Accossion NO: 10340 Top Hit Accession Accossion NO: 10340 Top Hit Accession Accossion NO: 10340 Top Hit Accession Accossion NO: 10340 Top Hit Accession Accossion NO: 10340 Top Hit Accession Accossion NO: 10340 Top Hit Accession Accossion NO: 10340 Top Hit Accession Accossion NO: 10340 Top Hit Accession Accossion NO: 10340 Top Hit Accession Accossion NO: 10340 Top Hit Accession Accossion NO: 10340 Top Hit Accession Accossion NO: 10340 Top Hit Accession Accossion Accossion NO: 10340 Top Hit Accession Accossion NO: 10340 Top Hit Accossion Accossion NO: 10340 Top Hit Accossion Accossion Accossion Accossion NO: 10440 Top Hit Accossion Accossio	Top Hit Descriptor	Homo sapiens IgG Fc binding protein (FC(GAMMA)BP) mRNA	Homo sapiens IgG Fc binding protein (FC(GAMMA)BP) mRNA	Homo sapiens IgG Fc binding protein (FC(GAMMA)BP) mRNA	Homo sapiens IgG Fc binding protein (FC(GAMMA)BP) mRNA	Homo sapiens IgG Fc binding protein (FC(GAMMA)BP) mRNA	Homo sapiens IgG Fc binding protein (FC(GAMMA)BP) mRNA	H.sapiens gene for RNA pol II largest subunit; exons 23-29	H.saplens gene for RNA pol II largest subunit, exons 23-29	Homo sapiens ribosomal protein L19 (RPL19) mRNA	yg09a02.r1 Scares infant brain 1NIB Homo scipiens cDNA clone IMAGE:31652 5'	Homo sapiens ribosomal protein S5 (RPS5) mRNA	Homo sapiens mRNA for KIAA1019 protein, partial cds	Homo sapiens SON DNA binding protein (SON) mRNA	Homo sapiens SON DNA binding protein (SON) mRNA	Mus musculus truncated SON protein (Son) mRNA, complete cds	Homo sapiens interferon gamma receptor 1 (IFNGR1) mRNA	EST27054 Cerebellum II Homo sapiens cDNA 5' end	601111520F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3352348 5'	Homo sapiens 5-hydroxytryptamine (serotonin) receptor 1B (HTR1B) mRNA	Homo sapiens 5-hydroxydryptamine (serotonin) receptor 1B (HTR1B) mRNA	Homo sapiens keratin 18 (KRT18) mRNA	Homo sapiens keratin 18 (KRT18) mRNA	Homo sapiens chromosome 21 segment HS21C046	Homo sapiens chromosome 21 segment HS21C046	Homo sapiens chromosome 21 segment HS21C046	Homo sapiens mRNA for KIAA1209 protein, partial cds	AU132898 NT2RP4 Hamo saplens cDNA clone NT2RP4000837 5	601274951F1 NIH_MGC_20 Homo saplens cDNA clone IMAGE:3615756 5'	PM0-DT0065-130400-002-c06 DT0065 Horniy sapiens cDNA	Novel human gene mapping to chomosome 1	Homo sapiens PC326 protein (PC326), mRNA	Homo sapiens chromosome 21 segment HS21C010	QV2-BT0635-160400-142-h05 BT0635 Homo saplens cDNA	601764858F1 NIH_MGC_53 Homo saplens cDNA clone IMAGE:3996998 5'
Expn SEQ ID ID NO: Check Similar Signal Most Similar Value Top Hit Acession Value Top Hit Acession No: Top Hit Acession Value Top Hit Acession No: Most Similar Value No. No. No. No. No. Most Similar No. No. No. No. No. No. No. No. No. No. No. No. No. No. No. No. A503880 No. No. A503880 No. No. A503880	Top Hit Database Source	TN	NT	NT	FN	N F	NT	NT	NT	NT	EST_HUMAN	NT	LN	TN	F	۲	NT	EST_HUMAN	EST_HUMAN	NT	NT	NT	NT	NT	NT	NT	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	TN	TN		EST_HUMAN
Exan Dor. 10338 ORF SEQ Expression Signal ID NO: Signal ID NO: Signal ID NO: Signal ID NO: 10338 Expression Signal ID NO: Dor. Dor. Dor. Dor. Dor. Dor. Dor. Dor.	Top Hit Acesston No.	4503680	4503680	4503680	4503680	4503680	4503680			ľ	795.1	4506728	128942.1	4507152		193607.1	4557879	124262.1	254447.1	4504532	4504532	4557887		63246.2	163249.2	63246.2		AU132898.1	3E385144.1		117233.1	8923955	63210.2		3F028005.1
Exan SEQ ID ID NO: Signa NO: NO: Signa NO: NO: NO: NO: NO: NO: NO: NO: NO: NO:		0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00)	0.0E+00)	0.0E+00	0.0E+00.	0.0E+00	0.05+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00)	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00
SEQ 10 NO: 10338 10338 10338 10338 10338 10338 10338 10338 10338 10338 10338 10338 10338 10338 10338 10338 10338 10338 10338 10404 10408 1	Expression Signal	2.04	1.17	1.64	1.64	2.43	6.0	0.84	0.84	43.09	1.31	2.61	2.42	4.7	4.7	3.51	2.01	0.98	0.91	3.15	3.15	1.27	1.27	2.26	7.05	7.05	2.59	1.64	2.17	1.05	1.07	1.64	3.91	1.97	1.13
	ORF SEQ ID NO:	20161	20162	20163	20164	20165	20166	20167	20168		19772		20190	20191	20192	20193	20206			20220	20221	20228	20229	20235	20236	20237	20242	20244	20250	20251	20253	20254	20263	20267	20273
SEQ D NO. O NO. O SEQ D	SEQ ID	10337	10338	10339	10339				H		1866	10367	10368	10369	10369	10370	10382	10387	10388	10404	10404	10408	10408	10419	10420	10420	10428	10430	10438	12668	10440	10441	10450	12669	10462
	Probe SEQ ID NO:	391	392	333	383	394	395	396	396	400	414	422	423	424	424	425	438	£	444	460	460	465	465	475	478	476	486	487	495	496	498	499	803	515	620

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Hamo saplens ubiquind-cytochrome c reductase, Rieske iron-sulfur polypeptide 1 (UQCRFS1), nuclear gene zh51b04.r1 Soares_fetal_liver_spleen_1NRLS_S1 Homo saplens cDNA clone IMAGE:415567 5' similar to gb:A21187 ALPHA-2-MACROGLOBULIN PRECURSOR (HUMAN); zh61b04,r1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:415567 5' similar to Homo saplens X-linked anhidrollic ectodermal dysplasia protein gene (EDA), exon 2 and flanking repeat Homo sapiens guanine nucleotide binding protein (G protein), elpha 11 (Gq class) (GNA11) mRNA Homo sapiens guanine nucleotide binding protein (G protein), alpha 11 (Gq class) (GNA11) mRNA UI-H-BI1-ecb-h-04-0-UI.s1 NCI_CGAP_Sub3 Homo sepiens cDNA clone IMAGE:2713951 3' Homo sapiens transcription elongation factor B (SIII), polypeptide 1-like (TCEB1L) mRNA Homo sapiens glutamate receptor, lonotropic, N-methyl D-aspartate 2B (GRIN2B) mRNA Homo sapiens CCAAT-box-binding transcription factor (CBF2) mRNA Homo sapiens sodium/calcium exchanger isoform NaCa3 (NCX1) mRNA, complete cds Homo sapiens sodium/calcium exchanger isoform NaCa3 (NCX1) mRNA, complete cds Homo saplens Smad- and Olf-Interacting zinc finger protein mRNA, partial cds Homo saplens Smad- and Olf-Interacting zinc finger protein mRNA, partial cds zt60c07.r1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:726732 5 601822627F1 NIH_MGC_75 Homo saplens cDNA clone IMAGE:4045447 5 Homo sapiens low density lipoprotein-related protein 2 (LRP2), mRNA Homo sapiens low density lipoprotein-related protein 2 (LRP2), mRNA Homo sapiens low density lipoprotein-related protein 2 (LRP2), mRNA Homo sapiens low density lipoprotein-related protein 2 (LRP2), mRNA Homo sapiens low density lipoprotein-related protein 2 (LRP2), mRNA Homo sapians acetyl-Coenzyme A carboxylase beta (ACACB), mRNA gb:A21187 ALPHA-2-MACROGLOBULIN PRECURSOR (HUMAN) Top Hit Descriptor Homo saplens NOD1 protein (NOD1) gene, exons 1, 2, and 3 Homo saplens novel SH2-containing protein 3 (NSP3) mRNA Human neutral amino acid transporter (ASCT1) gene, exon 8 Homo sapiens mRNA for KIAA1476 protein, partial cds Homo sapiens protein kinase, X-linked (PRKX) mRNA Homo sapiens RGH1 gene, retrovirus-like element -luman apolipoprotein A-I (ApoA-I) gene, excit Homo sapiens RGH2 gene, retrovirus-like encoding mitochondrial protain, mRNA Single Exon Probes Expressed in Heart regions Top Hit Database Source HUMAN HUMAN HUMAN **EST HUMAN** EST_HUMAN NT EST EST EST 4504036 NT 4504036 NT 눋 ż 눋 눋 6806918 NT 5031624 NT 닐눋 5174742|NT 4501854 NT 뉟 6806918 NT 6806918 NT 6806918 NT 4885526 NT 6006003 NT 6806918 h Fop Hit Acession 135324.1 0.0E+00 AF003528.1 0.0E+00 AW135324.1 0.0E+00 AF221712.1 0.0E+00 AF108389.1 AA399486.1 0.0E+00 AF108389.1 0.0E+00 482 AB040909.1 BF104898.1 F221712.1 ġ AF149773. 0.0E+00 W78811.1 W78811.1 0.0E+00 D10083.1 0.0E+00 D11078.1 0.0E+00/ 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 (Top) Hit BLAST E Value 1.16 2.18 2.18 1.79 0.83 3.98 3.98 3.15 5.28 1.46 1.05 1.38 98.0 222 0.93 1.34 3.17 3.28 2.89 9. 1.41 3.98 6.37 Expression Signal 20343 20351 20354 20283 20318 20355 20375 20395 20398 ORF SEQ 20284 20295 20357 20376 20388 20391 20399 20285 20372 ÖΝΩ SEQ ID 10468 10472 10472 10477 10485 10510 10529 10534 10546 10547 10548 10578 10579 10471 10495 10522 10543 10547 10556 10564 10564 10574 10583 10588 10548 10560 10567 ÿ Probe SEQ ID 526 529 530 536 644 571 587 583 588 598 598 607 610 612 619 830 639 946 846 846 652 630 554 611 g 627 ÿ

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Top Hit Descriptor	Homo sapiens protein kinase, X-linked (PRKX) mRNA	Human endogenous retrovirus pHE.1 (ERV9)	Homo saplens high-mobility group (nonhistone chromosomal) protein 1 (HMG1) mRNA	Homo sapiens mRNA for KIAA1089 protein, partial cds	Homo sapiens similar to rat integral membrane glycoprotein POM121 (POM121L1), mRNA	np49d01.s1 NCI_CGAP_Br1.1 Homo sepieris cDNA clone IMAGE:1129633 3' similar to gb:X57352 INTERFERON-INDUCIBLE PROTEIN 1-8U (HUMAN);	Human von Willebrand factor gene, exons 23 through 34	Human von Willebrand factor gene, exons 23 through 34	Homo sapiens TNF receptor-associated factor 1 (TRAF1) mRNA	Homo sapiens ALR-like protein mRNA, partial cds	Homo sapiens ALR-like protein mRNA, partial cds	Homo sapiens hypothetical protein FLJ21634 (FLJ21634), mRNA	TCAAP1D0779 Pediatric acute myslogenous leukemia cell (FAB M1) Baylor-HGSC project≂TCAA Homo saniens cDNA clone TCAAP0779	Homo saplens MHC class I antigen (HLA-G) mRNA, HLA-G1 allele, complete cds	Homo sapiens MHC class I antigen (HLA-G) mRNA, HLA-G1 allele, complete cds	Homo sapiens chloride channel CLC4 (CIC4) mRNA, complete cds	Human, plasminogen activator inhibitor-1 gene, exons 2 to 9	Human, plasminogen activator inhibitor-1 gene, exons 2 to 9	Homo sapiens mRNA for KIAA1339 protein, partial cds	Homo sapiens zinc finger protein 212 (ZNF212), mRNA	Homo sapiens mRNA for repressor protein, partial cds	601445647F1 NIH_MGC_65 Homo sapiens cDNA done IMAGE:3849803 5'	yj69g08.r1 Soares breast 2NbHBst Homo sapiens cDNA clone IMAGE:154046 5'	Homo sapiens splicing factor 3a, subunit 1, 120kD (SF3A1), mRNA	Homo sapiens gene for AF-6, complete cds	Homo sapiens KIAA0170 gene product (KIA40170), mRNA	Human mRNA for KIAA0184 gene, partial cds	Human mRNA for KIAA0184 gene, partial cd's	H.saptens mRNA for Interferon alpha/beta receptor (long form)	Homo sapiens mRNA for KIAA0910 protein, partial cds	Homo sapiens mRNA for KIAA0910 protein, partial cds	Homo saplens pericentrin (PCNT) mRNA
Top Hit Database Source	NT	NT	N	Z	Ę	EST HUMAN	NT L	NT	FX	LN TA	둗	FZ	EST HIMAN	L	N FA	LN	LN	NT	FN	N FA	FZ	EST_HUMAN	EST_HUMAN	LN	LN	TN	LZ LZ	NT	IN	NT	NT	ΝΤ
Top Hit Acession No.	4826947 NT	K57147.1	4504424 NT	AB029012.1	7657468 NT	AA614537.1		M60675.1	5032192 NT	AF264750.1	AF264750.1	11545800 NT	RF241577 1	Ī		AF170492.1	J03764.1	J03764.1	AB037760.1	6912749 NT	D30612.1	BE869735.1		5032086 NT	AB011399.1	7661965 NT	D80006.1	D80006.1		AB020717.1	AB020717.1	5174478 NT
Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00/	0.0E+00		0.0E+00 /		0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.05+00.0	_	+	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00/	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00
Expression Signal	3.98	96.0	4.56	4.35	2.03	19.46	7.66	7.68	1.45	3.89	3.89	87.8	1.7	1.12	1.12	2.4	1.55	1.55	0.78	1.12	0.81	2.17	3.38	2.4	1.58	2.97	1.17	1.17	2.64	2.37	2.37	6.84
ORF SEQ ID NO:	20405		20419	20423		20450	20454	20455	20464	20470	20471	20474	20481	20505	20506	20507	20510	20511	20512	20513	20514	20515	20519	20520	20529	20533	20544	20545	20549	20553	20554	20560
Exon SEQ ID NO:	10588	12672	10801	10608	10615	10625	10629	10629	10639	10644	10844	10646	10851	10670	10670	10871	10674	10674	10876	10677	12676	10678	10682	10683	10692	10698	10705	10705	10710	10714	10714	10718
Probe SEQ ID NO:	652	828	687	672	682	692	969	969	708	712	712	714	719	739	739	740	743	743	745	748	747	748	752	753	762	765	775	775	780	784	784	789

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Homo sapiens potassium voltage-gated channel, isk-related family, member 1 (KCNE1) mRNA Homo sapiens GA-binding protein transcription factor, alpha subunit (60kD) (GABPA), mRNA Homo sapiens T-cell lymphoma Invasion and metastasis 1 (TIAM1) mRNA Homo sapiens hormonally upregulated neu tumor-associated kinase (HUNK), mRNA Homo sapiens hormonally upregulated neu tumor-associated kinase (HUNK), mRNA Homo sapiens hormonally upregulated neu turnor-associated kinase (HUNK), mRNA Homo sapiens hormonally upregulated neu tumor-associated kinase (HUNK), mRNA Homo sapiens hormonally upregulated neu tumor-associated kinase (HUNK), mRNA Homo sapiens laminin receptor 1 (67kD, ribosomal protein SA) (LAMR1), mRNA (67kD, ribosomal protein SA) (LAMR1), mRNA Homo saplens sodium/myo-Inosital cotransporter (SLC5A3) gene, complete cds Homo sapiens hormonally upregulated neu tunior-associated kinase (HUNK), Homo sapiens serine-threonine protein kinase (MNBH) mRNA, complete cds 602085579F1 NIH_MGC_83 Homo sapiens cDNA clane IMAGE:4249915 5' Homo sapiens serine-threonine protein kinase (MNBH) mRNA, complete cds protein C inhibitor [human, leukocytes, Genomic, 1216 nt, segment 2 of 5] protein C inhibitor [human, leukocytes, Genomic, 1216 nt, segment 2 of 5] protein C Inhibitor [human, leukocytes, Genomic, 1216 nt, segment 2 of 5] Homo sapiens T-cell lymphoma invasion and nietastasis 1 (TIAM1) mRNA Homo sapiens T-cell lymphoma invasion and metastasis 1 (TIAM1) mRNA nj66d07.s1 NCI_CGAP_Pr10 Homo sapiens cDNA clone IMAGE:997453 ni66d07.s1 NCI CGAP Pr10 Homo sapiens DNA clone IMAGE:997453 Homo sapiens alpha-1-antichymotrypsin precursor, mRNA, partial cds Top Hit Descriptor QV0-BT0703-280400-211-g11 BT0703 Homo saplens cDNA QV0-BT0703-280400-211-g11 BT0703 Homis sapiens cDNA Homo sapiens mRNA for KIAA1019 protein, partial cds Homo saplens SON DNA binding protein (SON) mRNA Homo saplens mRNA for KIAA0910 protein, partial cds Homo sapiens mRNA for KIAA1019 protein, partial cds Homo sapiens mRNA for KIAA 1019 protein, partial cds Homo sapiens mRNA for KIAA0910 protein, partial cds Homo sepiens chromosome 21 segment HS21C003 Homo sapiens ribosomal protein S5 (RPS5) mRNA Homo sepiens chromosome 21 segment HS21C00 Homo sepiens laminin receptor 1 Single Exon Probes Expressed in Heart EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN HUMAN Top Hit Database Source 뉟 4507500 NT 7657213 NT LΝ 눋 4503854 NT 4507500 NT 눌 4506728 NT 7657213 NT 7657213 NT 4557686 NT 4507152 NT 7657213 NT 7657213 7657213 4504958 4507500 4504958 Top Hit Acession 0.0E+00 45077 0.0E+00 AB028942.1 0.0E+00 45067 0.0E+00 AB020717.1 0.0E+00 AB020717.1 0.0E+00 AL163203.2 AL163203.2 0.0E+00|AF108830.1 AF027153.1 AB028942.1 AA533272.1 BF677694.1 0.0E+00 S69364.1 0.0E+00 S69364.1 AB028942.1 AF089747 ģ 0.0E+00 0.0E+00/ 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0€+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 (Top) Hit BLAST E Jost Similar 216 3.93 0.87 1.85 1.85 7.45 3.29 0.99 0.99 7.39 42 . 15 7.38 2.02 1.97 7.08 1.57 3.37 3.37 1.97 Expression Signal 9 20874 20676 20618 20622 20626 20650 20614 20615 20617 20620 20665 20590 20613 20621 20623 20824 20591 20601 ORF SEQ ÖNQ 10763 10825 10825 10828 10829 10829 10744 10744 10770 10776 10789 10806 10816 10829 10752 10759 10763 10766 10770 10771 10772 10776 10777 10719 10764 10737 10739 SEQID 1077 8880 880 800 800 804 905 905 808 838 837 838 839 843 843 44 8 44 44 845 850 849 850 850 873 SEQ ID 816 816 780 807 808 810 825 825 832 22

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Table 4
Single Exon Probes Expressed in Heart

		_	_	_	_			_		-							_	_		ŋ	-	, ,				-17		1-1		<u></u>	f15	1.11	er pr=q 11
	Top Hit Descriptor	Homo sapiens kallistatin (PI4) gene, exons 1-4, complete cds	Homo saplens of cardiac alpha-myosin heavy chain gene	Homo sapiens of cardiac alpha-myosin heavy chain gene	Homo sapiens of cardiac alpha-myosin heavy chain gene	Homo saplens of cardlac alpha-myosin heavy chain gene	Human ras inhibitor mRNA, 3' end	Human ras inhibitor mRNA, 3' end	Human ras inhibitor mRNA, 3' end	Homo sapiens thyrotrophic embryonic factor (TEF), mRNA	Homo sapiens thyrotrophic embryonic factor (TEF), mRNA	os98e03.s1 NCI_CGAP_GC3 Homo saplens cDNA clone IMAGE:1613404 3'	os98e03.s1 NCI_CGAP_GC3 Homo sapiens cDNA clone IMAGE:1613404 3'	Homo sapiens KIAA0929 protein Msx2 Interacting nuclear target (MINT) homolog (KIAA0929), mRNA	Homo sapiens mRNA for PSP24, complete cds	PM2-GN0014-050900-001-f02 GN0014 Homo(sapiens cDNA	PM2-GN0014-050900-001-f02 GN0014 Homo saplens cDNA	PM2-GN0014-050900-001-f02 GN0014 Homo/sapiens cDNA	Homo sapiens partial c-fgr gene, exons 2 and 3	Homo sapiens partial c-fgr gene, exons 2 and 3.	Homo sapiens chromodomain protein, Y chromosome-like (CDYL) mRNA	Human beta-tubulin (TUB4q) gene, complete cds	Human beta-tubulin (TUB4q) gene, complete cds	Human beta-tubulin (TUB4q) gene, complete cds	Homo sapiens 8q22.1 region and MTG8 (CBF/\2T1) gene, partial cds	Homo saplens 8q22.1 region and MTG8 (CBFA2T1) gene, partial cds	Homo sapiens 14q32 Jagged2 gene, complete cds; and unknown gene	Homo sapiens 14q32 Jagged2 gene, complete cds; and unknown gene	Homo sapiens 14q32 Jagged2 gene, complete cds; and unknown gene	Homo sapiens 14q32 Jagged2 gene, complete cds; and unknown gene	Homo sapiens DKFZP586M0122 protein (DKFZP586M0122), mRNA	Homo sapiens inner membrane protein, mitochondrial (mitofilin) (IMMT), mRNA	aa88g07.s1 Stratagene fetal retina 937202 Homo sapiens cDNA clone IMAGE:838236 3' similar to SW:PRS8_HUMAN P47210 26S PROTEASE REGULATORY SUBUNIT 8;
, _	Top Hit Database Source	뉟	F	N	뉟	Z	Ā	뉟	ΙN	Z	Z	EST_HUMAN	EST_HUMAN	۲	NT.	EST HUMAN	EST HUMAN	EST_HUMAN	LN	NT	ΙN	NT	NT	NT	NT	NT	N	N	٦	۲	F	N	EST_HUMAN
	Top Hit Acessian No.	1,28101.1	220656.1	Z20656.1	Z20656.1	Z20656.1	0.0E+00 M37190.1	0.0E+00 M37190.1	M37190.1	4507430 NT	4507430 NT	AJ001948.1	0.0E+00 AI001948.1	7657266 NT	0.0E+00 AB030566.1	0.0E+00 BF366974.1	0.0E+00 BF366974.1	0.0E+00 BF366974.1	X52207.1	X52207.1	4757969 NT	U83668.1	J83668.1	J83668.1	0.0E+00 AF198490.1	0.0E+00 AF198490.1	0.0E+00 AF111170.3	0.0E+00 AF111170.3	0.0E+00 AF111170.3	0.0E+00 AF111170.3	7661685 NT	5803114 NT	0.0E+00 AA458680.1
	Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00 Z	0.0E+00 Z20656.1	0.0E+00 Z20656.1	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 A	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	O.0E+00	0.0E+00	0.0E+00	0.0E+00 U83668.1	0.0E+00 U83668.1	0.0E+00 U83668.1	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00
	Expression Signel	2.14	105.13	105.13	167.64	167.64	36.3	13,25	63.71	1.9	1.9	2.48	2.46	7.21	2.52	4.64	4. 8.	4.64	1.27	. 1.27	1.25	1.05	7.48	69.69	1.99	3.49	0.84	1.43	1.81	223	2.28	2.68	1.94
	ORF SEQ ID NO:		20680								20710	20717	20718	20720	20730	20736		20738	20739	20740			20756	20756			20761	20761	20761		20765		
	Exen SEQ ID NO:	10830	10833	10833	10834	10834	10860	10861	10862	10863	10863	12681	12681	10872	10882	10890	10890	10890	10891	10891	10900	10910	10911	10911	10914	10914	10917	10917	10917	10918	10921	10925	10927
	Probe SEQ ID NO:	908	808	608	910	910	935	936	937	938	838	948	946	848	626	296	296	987	898	988	977	888	686	990	993	994	997	866	666	1000	1003	1007	1009

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EST51i24 WATM1 Homo saplens cDNA clone 51i24 similar to DNA-DIRECTED RNA POLYMERASE II EST51124 WATM1 Homo sapiens cDNA clone 51124 similar to DNA-DIRECTED RNA POLYMERASE II Homo sapiens glutamate decarboxylase 1 (brain, 67kD) (GAD1), transcript variant GAD25, mRNA Homo sapiens glutamate decarboxylase 1 (brain, 67kD) (GAD1), transcript variant GAD25, mRNA NbHPU Homo sapiens cDNA clone IMAGE:1697011 3' domo sapiens similar to rat integral membrane glycoprotein POM121 (POM121L1), mRNA Homo sapiens similar to rat integral membrane glycoprotein POM121 (POM121L1), mRNA Homo sapiens TRAF family member-associated NFKB activator (TANK) mRNA Homo sapiens TRAF family member-associated NFKB activator (TANK) mRNA Homo sapiens potassium channel, subfamily K, member 9 (KCNK9), mRNA Homo sapiens heat shock 70kD protein 9B (mortalin-2) (HSPA9B) mRNA Homo saplens chromosome 12 open reading frame 3 (C12ORF3), mRNA fomo sapiens potassium channel, subfamily KI, member 9 (KCNK9) Homo sapiens cadherin 6, K-cadherin (fetal kidney) (CDH6) mRNA Homo saplens Npw38-binding protein NpwBP (LOC51729), mRNA Homo sapiens cadhertn 6, K-cadherin (fetal kidney) (CDH6) mRNA Homo sapiens hypothetical protein FLJ20695 (FLJ20695), mRNA Homo sapiens hypothetical protein FLJ11196 (FLJ11196), mRNA Homo saplens hypothetical protein FLJ20695 (FLJ20695), mRNA mRNA Homo sapiens hypothetical protein FLJ20309 (FLJ20309), mRNA Top Hit Descriptor VRO-BN0115-200300-003-h08 BN0115 Homo sapiens cDNA Homo sapiens alkylation repair, alkB homolog (ABH), mRNA Homo sapiens mRNA for alpha-tubulin 8 (TUBA8 gene) Homo sapiens hypothetical protein FLJ20080 (FUJ20080) Homo sapiens Death associated protein 3 (DAP3) mRNA Homo sapiens ribosomal protein S27a (RPS27A) mRNA Homo sapiens mRNA for KIAA1414 protein, partial cds tomo sapiens mRNA for KIAA0903 protein, partial cds lomo saplens protein kinase, X-linked (PRKX)|mRNA Homo sapiens protein kinase, X-linked (PRKX) mRNA Homo saplens DNA for Human P2XM, complete cds fomo sapiens DNA for Human P2XM, complete cds alignment Ser and Pro with BLASTx or p) (alignment Ser and Pro with BLASTx or p qb22d10.x1 Soares_pregnant_uterus EST_HUMAN HCMAN Top Hit Database HUMAN Source EST 22.1 NT 8923087 NT 5174384 NT EST 7706134 NT 4826947 NT 4759249 NT 4759249 NT 8922933 NT 4758569|NT 4826672 NT 8923624 NT 4758117 NT 4826947 NT 4506712 NT 8923290 NT IN 9966844 NT 7305076 NT 4826672 NT 7706134 NT 7657468 NT 7857488 NT 7706500 NT 7305076 NT Top Hit Acession 0.0E+00 8923 0.0E+00 AJ245922.1 0.0E+00 AB002059.1 0.0E+00 AB020710.1 AB002059.1 AB037835.1 ģ 0.0E+00 N43182.1 N43182.1 0.0E+00 0.011-00 0.0E+00 0.0E+00 (Top) Hit BLAST E 0.0E+00 dost Similar 38.24 3.28 3.58 3.79 0.86 <u>\$</u> 0.97 0.97 4.44 1.88 1.88 2.52 2.52 10.18 12.02 2.03 0.951.13 2.31 2.31 1.59 5.7 2.03 Expression Signal 20808 20812 20813 20862 20872 20873 20876 ORF SEQ ID NO: 20775 20878 20879 20880 20774 20814 20861 20874 20884 20792 20827 20887 20777 SEQ ID 10966 10966 10930 10930 10949 10975 10982 11019 11045 10970 10970 11019 11038 11039 11054 10931 10971 10973 11031 11036 192 11031 1065 11067 Š ÿ Probe SEQ ID 1049 1049 1116 1013 1053 1053 1056 1058 1068 1103 1116 1119 1123 1124 1128 1131 146 1152 1154 1912 1012 1013 1054 1103 1124 80 13 1129 1631 ÿ

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		_	_	_	_	_	_		,	_	~			_	_	_	_	_	_		***	-		***					٠,٠		4	11 11 11	11 11 11 11
	Top Hit Descriptor	Homo sapiens keretin 18 (KRT18) mRNA	Homo sapiens mutt. (E. coli) homolog 3 (MLH3), mRNA	Homo sapiens ALR-like protein mRNA, partial cds	Homo sapiens ALR-like protein mRNA, partial cds	Homo sapiens ALR-like protein mRNA, partial ¢ds	Homo sapiens ALR-like protein mRNA, partial ¢ds	Homo sapiens chromosome 3 subtelameric region	Homo saplens chondroitin suffate proteoglycan 4 (melanoma-associated) (CSPG4), mRNA	Homo sapiens NF2 gene	Homo sapiens ribosomal protein S2 (RPS2) mRNA	Homo saplens Williams-Beuren syndrome deletion transcript 9 (WBSCR9) mRNA, complete cds	Homo sapiens mRNA for KIAA1507 protein, pertial cds	Homo sapiens mRNA for KIAA1507 protein, pertial cds	Homo sapiens Wolfram syndrome (WFS) mRNA	Homo sapiens Wolfram syndrome (WFS) mRNA	Homo saplens Wolfram syndrome (WFS) mRNA	Homo sapiens protein phosphatase 2A BR gamma subunit gene, exon 5	Homo saplens rhabdoid tumor deletion region protein 1 (RTDR1), mRNA	Homo saplens rhabdoid tumor deletion region protein 1 (RTDR1), mRNA	Homo sapiens ring finger protein 9 (RNF9), mRNA	Homo saplens zinc finger protein 173 (ZNF173) mRNA	Homo saplens ring finger protein 9 (RNF9), mRNA	Homo sapiens mRNA for KIAA0577 protein, complete cds	Homo sapiens KIAA0170 gene product (KIAA0170), mRNA	Homo sapiens KIAA0170 gene product (KIAA0170), mRNA	Homo sapiens period (Drosophila) homolog 3 (PER3), mRNA	Homo sapiens period (Drosophila) homolog 3 (PER3), mRNA	Human endogenous retrovirus HERV-K10	Homo sapiens mRNA for Familial Cylindromatissis cyld gene	Homo sapiens partial TTN gene for titin	qg38b06.x1 Soares_testis_NHT Homo sapien's cDNA clone IMAGE:1837427 3' similar to WP:T27A1.5 CE14213 ;	RAN, member RAS oncogene familyHomo sapiens RAN, member RAS oncogene family (RAN), mRNA
,	Top Hit Database Source	눌	Z.	Z	N	FZ	LN FN	Z	닐	NT	NT	. IN	NT	NT	NT	NT	NT	M	NT	NT	NT	NT	NT	NT	NT	NT	NT	NT	IN	IN	IN	EST_HUMAN	FZ.
	Top Hit Acessian No.	TN 4557887 NT	7657336 NT	AF264750.1	AF264750.1	AF264750.1	AF264750.1	AF109718.1	4503098 NT	Y18000.1	4506718 NT	AF084479.1		AB040940.1	5174748 NT	5174748 NT	5174748 NT	AF096156.1	7657529 NT	7657529 NT	5803146 NT	4508004 NT	3146	AB011149.1	7681965 NT	7661965 NT	8567387 NT	8567387 NT	M14123.1	AJ250014.1	AJ277892.1	AI208756.1	6042206 NT
	Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00)	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00
	Expression Signal	1.02	1.13	1.14	1.14	1.31	96'0	3.62	1.3	1.48	45.69	3.66	1.71	1.71	6.42	6.42	6.42	. 2.6	1.1	1.1	1.71	0,82	1.07	4.1	1.06	4.64	4.1	4.1	1.35	0.89	9.13	0.98	8.18
	ORF SEQ ID NO:	20920		20962	20963		20965	20988	20989							21037	21038								21062		21084		21078	21147	21156	21160	21161
	Exon SEQ ID NO:	11074	11101	11116	11116		12687	11134	11135							11186				i							11210		11221	11291	11298	11301	11302
	Probe SEQ ID NO:	1161	1191	1206	1206	1207	1208	1226	1227	1245	1253	1280	1266	1266	1278	1278	1278	1279	1289	1289	1295	1298	1298	1300	1301	1302	1303	1333	1315	1386	1393	1398	1397

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	Top Hit Descriptor Top Hit Descriptor Source	IT Homo sapiens proprotein convertase subtilisin/kaxin type 2 (PCSK2) mRNA	IT Homo saplens proprotein convertase subtilisin/lexin type 2 (PCSK2) mRNA					IT Homo saplens titn (TTN) mRNA	IT Homo saplens titn (TTN) mRNA	NT Human nebulin mRNA, partial cds	NT Human nebulin mRNA, partial cds		NT Novel human gene mapping to chomosome 1	NT Human mRNA for KIAA0240 gene, partial cds,	iT Homo sapiens calcineurin binding protein 1 (KIAA0330), mRNA	IT Homo sapiens KIAA0170 gene product (KIAA0170), mRNA	IT Homo sapiens KIAA0170 gene product (KIAA0170), mRNA	NT Homo sepiens RFB30 gene for RING finger protein		NT Human von Wilebrand factor pseudogene corresponding to exons 23 through 34		EST_HUMAN as34803.11 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:815116 5'		NT Cercopithecus aethiops cyclophilin A mRNA, complete ods	NT Bovine mRNA for neurocalcin						IT Homo sapiens TNF-inducible protein CG12-1 (CG12-1), mRNA	NT Human transglutaminase mRNA, complete cds			IT Homo sapiens ribosomal protein L5 (RPL5) mRNA
-	Top Hit Acession No.	4505646 NT	4505646 NT	7705565 NT	7705565 NT	AJ238093.1	AF038280.1 N	4507720 NT	4507720 NT	U35637.1	U35637.1 N		AL137764.1	D87077.1	6912457 NT	7661965 NT	7661965 NT	Y07829.2		M60676.1	7706434 NT		AF023860.1	AF023860.1 N	D10884.1		U78027.1	4505404 NT	4505404 NT	7662405 NT	7656972 NT	M98478.1 N	4507720 NT	4507720 NT	4506654 NT
	Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00
	Expression Signal	4.4	1.4	2.54	2.54	5.25	3.51	9.7	9.7	1.02	1.02	3.05	1.03	122	4.97	1.51	1.51	0.97	3.65	3.65	1.32	0.95	11.95	11.95	16.0		2.83	3.9	3.9	3.12	8.41	5.02	5.76	5.75	10.12
	ORF SEQ ID NO:	21173	21174	21177	21178	21180	21192	21203	21204	21208	21209	21214	21215	21220	21223	21225	21226		21231	21232	21259	21273	21276	21277	21280			21283	21284	21285		21289		21292	
	Exan SEQ ID NO:	11312	11312	11314	11314	11317	11327	11337	11337	11342	11342	11350	11352	11356	11359	11361	11361	11362	11367	11367	L	11414	11420	11420	11422	1	ı	11425	11425	11426	11427	11432	11435	11435	12697
	Probe SEQ ID NO:	1407	1407	1409	1409	1412	1421	1432	1432	1437	1437	1445	1447	1461	1454	1458	1456	1457	1462	1462	1495	1509	1515	1515	1517		1519	1520	1520	1521	1522	1527	1530	1630	1531

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Тор Hit Descriptor
1632	11436	21293	11.11	00+30'0	M14199.1	NT	Human laminin receptor (2H5 epitope) mRNA, 5' end
1541	11448	21306	5.81	00+30'0	4507720 NT	NT	Homo saplens titin (TTN) mRNA
1541	11448	21307	5.81	0.0E+00	4507720 NT	LN	Homo sapiens titin (TTN) mRNA
1543		21308	12.91	0.0E+00	4503098 NT	LN	Homo sapiens chondroitin sulfate protecglycan 4 (melanoma-associated) (CSPG4), mRNA
1662	_	21325	1.66	0.0E+00	Z83738.1	NT	H.saplens hH2B/e gene
1583	11468	21328	1.38	0.0E+00	5921460 NT	Z	Homo sapiens butyrophilin, subfamily 2, member A1 (BTN2A1), mRNA
1583	11468	21327	1.38	00+30'0	5921460 NT		Homo sapiens butyrophilin, subfamily 2, member A1 (BTN2A1), mRNA
1564	11469		5.5	0.0E+00			AV690831 GKC Homo sapiens cDNA clone GKCBOF02 5'
1564	11469	21329	5.5	0.0E+00	AV690831.1	EST_HUMAN	AV690831 GKC Homo sapiens cDNA clone GKCBOF02 5
1566	12698	21330	1.5	0.0E+00		NT	Homo sapiens mRNA for KIAA1472 protein, pertial cds
1570		21331	86.0	0.000	AF157476.1	NT	Homo sapiens DNA polymerase zeta catalytic subunit (REV3) mRNA, complete cds
1572	11476	21334	2.49	0.0E+00	7662183 NT	NT	Homo sapiens KIAA0569 gene product (KIAA0569), mRNA
1572	11476	21335	2.49	0.0E+00	7662183 NT	NT	Homo sapiens KIAA0569 gene product (KIAA0569), mRNA
1574	11478	21336	14.05	0.0E+00	5729876 NT	NT	Homo sapiens heat shock 70kD protein 10 (HSC71) (HSPA10), mRNA
1574	11478	21337	14.05	0.0E+00	5729876 NT	LN T	Homo sapiens heat shock 70kD protein 10 (HSC71) (HSPA10), mRNA
1576	11480	21339	0.94	0.0E+00	M91803.1	TN	Human sodium channel mRNA
1591	11495	21355	4.67	0.0E+00	H26973.1	EST_HUMAN	yo76c05.s1 Scares adult brain N2b4HB55Y Homo sapiens cDNA clone IMAGE:183848 31
1602	11507	21368	1.4	0.0E+00	AB046829.1	NT	Homo sapiens mRNA for KIAA1609 protein, partial cds
1602	11507	21369	1.4	0.0E+00	AB046829.1	NT	Homo sapiens mRNA for KIAA1609 protein, partial cds
1645	11549	21410	1.27	0.0E+00	AI768104.1	EST_HUMAN	wg81b07.x1 Soares_NSF_F8_9W_OT_PA_P.31 Homo sapiens cDNA clone IMAGE:2371477 3' similar to TR:Q62788 Q62788 CYS2HIS2 ZINC FINGËR PROTEIN ;
1848	١	21411	3.33	0.05+00	AF057177.1	F	Homo sapiens T-cell receptor gamma V1 gene region
1650	_		1.58	0.0E+00		NT	Human zinc-finger protein 7 (ZFP7) mRNA, complete cds
1650	<u></u>		1.56		M29580.1	NT	Human zinc-finger protein 7 (ZFP7) mRNA, complete cds
1652	11555	21418	1.22	0.0E+00	TN 288734	NT	Homo sapiens keratin 18 (KRT18) mRNA
1653	11556	21419	0.92			NT	Homo sapiens v-ets avian erythroblastosis virus E26 oncogene related (ERG), mRNA
1657	11559		1.11	0.0E+00	TN 0197554	IN	Homo saplens gamma-aminobutyric acid (GABA) A receptor, gamma 2 (GABRG2) mRNA
1659	11561	21426	2.99		H30132.1	EST_HUMAN	yo59e08.r1 Soares breast 3NbHBst Homo sapiens cDNA clone IMAGE:182246 5' similar to gb:M64099 GAMMA-GLUTAMYLTRANSPEPTIDASE 5 PRECURSOR (HUMAN);
1669			2.99	0.0E+00	H30132.1	EST_HUMAN	yo59e08.r1 Soares breast 3NbHBst Homo sapiens cDNA clone IMAGE:182246 5' similar to gb:M64099 GAMMA-GLUTAMYLTRANSPEPTIDASE 5 PRECURSOR (HUMAN);
1681	┖		1.21		280780.1	E	H.sapiens H2B/h gene
1991	11563		1.21	0.0E+00	Z80780.1	L'N	H.sapiens H2B/h gene
1664	丄		7.85	0.0E+00	5031748 NT	NT	Homo sapiens high-mobility group (nonhistone chromosomal) protein 17 (HMG17), mRNA
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PCT/US01/00666

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	Top Hit Descriptor	Homo sapiens FOXJ2 forkhead factor (LOC55810), mRNA	Human hepatocyte growth factor gene, exon 15	Human hepatocyte growth factor gene, exon 15	Homo saplens RNA binding motif protein, Y chromosome, family 1, member A1 (RBMY1A1) mRNA	Homo sapiens WAVE2 mRNA for WASP-family protein, complete cds	TCR zata [human, Genomic/mRNA, 365 nt, segment 1 of 8]	Homo sapiens NOD2 protein (NOD2), mRNA	Homo saplens SMCY (SMCY) gene, complete cds	Homo sapiens ribosomal protein S2 (RPS2) mRNA	Homo sapiens E1A binding protein p300 (EP300) mRNA	Homo sapiens E1A binding protein p300 (EP300) mRNA	Human CSF-1 receptor (FMS) gene, complete cds, and (SMF) gene, partial cds	zd66g09.r1 Soares, fetal, heart, NbHH19W Homo sapiens cDNA clone IMAGE:345664.5	Homo saplens nuclear autoantigenic sperm protein (histone-binding) (NASP) mRNA	Human ribosomal protein L21 mRNA, complete ods	Human mRNA for KIAA0333 gene, partial cds	Homo sapiens activating transcription factor 4 (tax-responsive enhancer element B67) (ATF4) mRNA	Homo sapiens activating transcription factor 4 (tax-responsive enhancer element B67) (ATF4) mRNA	Homo saplens activating transcription factor 4 (tax-responsive enhancer element B67) (ATE4) mRNA	Homo saplens Immunoglobin superfamily, member 3 (IGSF3) mRNA, and translated products	Homo seplens Immunoglobin superfamily, member 3 (IGSF3) mRNA, and translated products	Homo sapiens Retina-derived POU-domain factor-1 (RPF-1), mRNA	Homo sepiens Retine-derived POU-domain factor-1 (RPF-1), mRNA	nily, member 1 (KCNB1) mRNA	Homo sepiens potassium voltage-gated channel Shab-related subfamily member 1 (KCNR4) mRNA			saplens cDNA clone IMAGE:2722333 3'	
) -	Top Hit Database Source	₽N FN	N F	N	NT	NT	TN	FN	FZ	NT	TN	N	TN	EST HUMAN	NT	ΓN	NT	NT	TN	ZT	Į,	ΙΝ	۲	TN	LΝ	Ę	NT	NT	EST_HUMAN	EST_HUMAN
	Top Hit Acession No.	8923841 NT	M75980.1	M75980.1	4826973 NT	AB026542.1	S94400.	11545911 NT	AF273841.1	4506718 NT	4557556 NT	4557556 NT	J63963.1	W76571.1	4505332 NT	U14967.1	AB002331.1	4502264 NT	4502264 NT	4502264 NT	4504626 NT	4504626 NT	6005855 NT	6005855 NT	4826783 NT	4826783 NT	J07147.1	J07147.1	0.0E+00 AW207280.1	0.0E+00 AW207280.1
	Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 U	0.0E+00/	0.0E+00/
	Expression Signal	4.55	1.02	1.02	1.43	2.59	2.8	0.87	1.75	13.81	6.0	6.0	1.23	1.13	3.89	7.25	4.6	4.34	4.34	4.34	1.04	1.04	10.22	10.22	3.19	3.19	4.52	4.52	1.47	1.47
	ORF SEQ ID NO:	21442	21448	21449	21453	21460			21481			21518			21525		21541	21542	21543	21544	21558	21559	21566	21567	21576	21677	21578	21579	21584	21585
	Exan SEQ ID NO:	11574	11579	11579			11589	12702	11611	12703	11649	11649	11651		12704	11664	11666	11667	11667	11667	11680	11680	11691	11691	11701	11701	11702	11702	11706	11706
	Probe SEQ ID NO:	1672	1677	1877	1680	1685	1687	1699	1710	1745	1749	1749	1751	1764	1755	1765	1787	1768	1768	1768	1781	1781	1793	1793	1804	1804	1805	1805	1809	1809

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Homo saplens glutathione S-transferase thela ? (GSTT2) and glutathione S-transferase thela 1 (GSTT1) Homo sapiens DNA polymerase zeta catalytic subunit (REV3) mRNA, complete cds Homo sapiens RAD1 (S. pombe) homolog (RAD1) mRNA, and translated products (RAI)1) mRNA, and translated products 601179164F1 NIH_MGC_20 Homo saplens cDNA clone IMAGE:3547239 5 x69b01.x1 NCL_CGAP_Pan1 Homo sapiens cDNA clone IMAGE:2878913 3 20 Homo sapiens cDNA clone IMAGE:3547239 Human topolsomerase i pseudogene 1 Homo sapiens butyrophilin, subfamily 3, member A2 (BTN3A2), mRNA Homo saplens transforming growth factor, beta 3 (TGFB3), mRNA Homo sapiens transforming growth factor, beta 3 (TGFB3), mRNA d69b01.x1 NCI_CGAP_Pan1 Homo sapiens cDNA clone IMAGE.: Tap Hit Descriptor Homo sapiens calcineurin binding protein 1 (KIAA0330), mRNA Homo sapiens calcineurin binding protein 1 (KIAA0330), mRNA RC2-BN0126-200300-012-b04 BN0126 Homo sapiens cDNA Homo sapiens mRNA for KIAA0790 protein, partial cds Human TFEB protein mRNA, partial cds Homo saplens mRNA for KIAA0790 protein, partial cds Homo sapiens mRNA for KIAA1513 protein, partial cds H.sapiens genes for semenogelin I and semenogelin II H.sapiens genes for semenogelin I and semenogelin I Homo saplens histidine ammonia-lyase (HAL) mRNA Homo saplens histidine ammonia-lyase (HAL) mRNA Homo sapiens chromosome 21 segment HS21,C052 Hamo saplens SMCY (SMCY) gene, complete cds Homo saplens SMCY (SMCY) gene, complete cds Homo sapiens actinin, alpha 4 (ACTN4) mRNA Human transglutaminase mRNA, complete cds Human transglutaminase mRNA, complete cds Homo sapiens actinin, alpha 4 (ACTN4) mRNA Homo sapiens RAD1 (S. pombe) homolog (Human TFEB protein mRNA, partial cds Homo sapiens nebulin (NEB), mRNA Homo sapiens nebulin (NEB), mRNA genes, complete cds 601179164F1 NIH HUMAN EST_HUMAN HUMAN Top Hit Database Source HUMAN EST_HUMAN 닐닏 4506384 NT 눋 4507464 NT 눌 뉟 불불 4506384 NT Ξ 5901905 NT 4809282 NT 눋 ΝŢ 닐눋 4507464[NT 4809282|NT z 8400716 NT 4826638 NT 6912457 NT 8400716 4826638 Top Hit Acession 6912457 0.0E+00 AW193024.1 AW 193024.1 AF157476.1 AF240786.1 AL163252.2 0.0E+00 AB018333.1 0.0E+00 AB040946.1 BE277465.1 BE006292.1 0.0E+00 AF273841.1 0.0E+00 AF273841.1 ģ 0.0E+00 AB018333. 0.0E+00 M33782.1 0.0E+00 M98478.1 0.0E+00 M33782.1 247556.1 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 (Top) Hit BLAST E Value 0.0E+00 0.0E+00 dost Simila 2.08 2.08 0.99 3.52 3.52 4.06 4.06 2.28 8.13 0.92 228 9,0 1.97 1.97 8.13 12 14 2.01 29 1.33 1.33 8.4 0.92 0.88 Expression Signal 21603 21623 21662 21669 21716 21723 21604 21651 21652 21670 21685 21720 ORF SEQ 21661 21702 21724 ÖΝΩ SEQ ID 11748 12707 11808 11819 11777 12707 11808 11821 11821 11822 11822 11832 11838 11840 11840 11841 11843 11868 11777 11832 11838 11841 11843 ÿ Probe SEQ ID 1913 1881 832 1926 1926 1943 1943 1945 1946 1946 1948 1948 1966 1975 1852 1881 883 1890 1890 895 1895 898 903 906 1913 1927 1927 1937 1937

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Table 4
Single Exon Probes Expressed in Heart

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Тэр Hit Descriptor	601573895F1 NIH_MGC_9 Homo septens cDNA clone IMAGE:3835198 5'	601573895F1 NIH_MGC_9 Homo sapiens cDINA clone IMAGE:3835198 5'	Homo sapiens coagulaton factor IX (plasma thromboplastic component, Christmas disease, hemophilia B) (F9) mRNA	AU140831 PLACE4 Homo sapiens cDNA clone PLACE4000321 5'	Homo sapiens KIAA1114 protein (KIAA1114), inRNA	Homo saplens KIAA1114 protein (KIAA1114), mRNA	7B22E10 Chromosome 7 Fetal Brain cDNA Library Homo saplens cDNA clone 7B22E10	7B22E10 Chromosome 7 Fetal Brain cDNA Library Homo sapiens cDNA clone 7B22E10	Homo sapiens similar to rat integral membrane glycoprotein POM121 (POM121L1), mRNA	Homo sapiens phosphodiesterase 6A, cGMP-specific, rod, alpha (PDE6A), mRNA	HSC0IC021 normalized Infant brain cDNA Horno sapiens cDNA clone c-0ic02	qv80f08.x1 NCI_CGAP_Ut2 Homo saplens cDNA clone IMAGE:1988871 3' similar to contains Alu repetitive element	601485146F1 NIH_MGC_69 Homo saplens cDNA clone IMAGE:3887747 5	801902604F1 NIH_MGC_19 Homo sapiens cDNA clane IMAGE:4135320 5	601902604F1 NIH_MGC_19 Homo sapiens cDNA clane IMAGE:4135320 5'	RC3-CT0413-270700-022-d10 CT0413 Homo saplens cDNA	RC3-CT0413-270700-022-d10 CT0413 Homo sapiens cDNA	Human plasma membrane calcium ATPase isoform 2 (APT2B2) mRNA, comlete cds	Human plasma membrane calcium ATPase isoform 2 (APT2B2) mRNA, comiete cds	Homo sapiens GTP binding protein 1 (GTPBP1) mRNA	QV1-GN0085-140800-318-c10 GN0065 Homo sepiens cDNA	Homo sapiens X-linked juvernile retinoschisis p/otein (XLRS1) gene, exon 6 and complete cds	601672066F1 NIH_MGC_20 Homo sapiens cDNA clone IMAGE:3954785 5'	Homo sapiens flavin containing moncoxygenase 2 (FMO2) mRNA	Homo sapiens glutathione S-transferase theta 2 (GSTT2) and glutathione S-transferase theta 1 (GSTT1) genes, complete cds	IL3-CT0219-271099-022-G10 CT0219 Homo sapiens cDNA	QV-BT065-020399-092 BT065 Homo saplens cDNA	QV-BT065-020399-092 BT065 Homo saplens cDNA	Homo sapiens potassium large conductance calcium-activated channel, subfamily M, beta member 3-like (KCNMB3L), mRNA	Human DNA-binding protein mRNA, 3'end
Top Hit Database Source	EST_HUMAN	EST_HUMAN	FN	EST HUMAN	۲N	L	EST_HUMAN	EST_HUMAN	· LN	FZ	EST_HUMAN	EST HIMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	N	N	N N	EST_HUMAN	FX	EST_HUMAN	NT	F	EST HUMAN	EST HUMAN	EST HUMAN	FZ	LN
Top Hit Acession No.	3E743215.1	3E743215.1	4503648 NT	\U14083		7705565 NT	AA077589.1	4A077589.1	7657468 NT	4585863 NT	242399.1	A124247 1	3E877225.1	3F315325.1	BF315325.1	3E697125.1	0.0E+00 BE697125.1	.00620.1	.00820.1	4758489 NT	3E767964.1	AF018963.1	3F027562.1	4503756 NT	4F240786.1	0.0E+00 AW752708.1	A1904640.1	1904640.1	7657252 NT	14787.1
Most Similar (Top) Hit BLAST E Value	0.0E+00 B	0.0E+00	0.0E+00	0.0E+00/	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 Z	0.05+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	00E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00
Expression Signal	1.09	1.09	98.0	5.48	1.01	1.01	1.95	1.95	2.34	6.1	0.82	194	2.59	1.5	1.5	2.42	2.42	2.53	2.53	1.32	2.63	1.13	3.09	2	0.99	1.16	1.86	1.96	1.19	1.6
ORF SEQ ID NO:		21794		21798		21178	21798	21799			21802		21811	L	L			21826		21831				21855		21857				
Exon SEQ ID NO:		11903	11805	l	11314	11314	11908	11908	11910	11912	11913	11915	11920	11922	L	l.	11928	11931	ı	11936	11957	11958	11960		11983	1	!	1		П
Probe SEQ ID NO:	2011	2011	2013	2014	2015	2016	2017	2017	2018	2021	2022	2024	2028	2031	2031	2035	2035	2040	2040	2045	2067	2088	2070	2071	2073	2074	2076	2076	2112	2132

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Top Hit Descriptor	601122338F1 NIH_MGC_20 Homo capiens cDNA clone IMAGE:3346688 5'	Human mRNA for KIAA0244 gene, partial cds	AV738288 CB Homo sapiens cDNA clone CBNBDE08 5'	AV738288 CB Homo sapiens cDNA clone CBNBDE08 5'	0032e01.s1 NCI_CGAP_Lu5 Homo sapiens cl0NA clone IMAGE:1567896 3'	Human apolipoprotein B-100 (apoB) gene, exons 22 through 29	602014829F1 NCI_CGAP_Bm64 Homo sapiens cDNA clone IMAGE:4150734 5	601572186T1 NIH_MGC_55 Homo saplens cDNA clone IMAGE:3839012 3'	CM1-TN0141-250900-439-b08 TN0141 Homc saplens cDNA	CM1-TN0141-250900-439-b08 TN0141 Home saplens cDNA	601900261F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:4129622 5	bb84e02.yi NIH_MGC_10 Hamo sapiens cDNA clane IMAGE:3049082 5' similar to TR:Q15170 Q15170	TRANSCRIPTION FACTOR SHIRELATED PROTEIN;	zk53c07.s1 Scares_pregnant_uterus_NbHPUIHomo sapiens cDNA cione IMAGE:486540 3' similar to	gb:X85857_cds1 OLFACTORY RECEPTOR-LIKE PROTEIN HGMP07E (HUMAN);	2k53c07.s1 Soares pregnant uterus. NbHPU/Homo sapiens cDNA clone IMAGE:486540 3' similar to	gb:Xebsb/ cast OLFACTORY RECEPTOR-LIKE PROTEIN HGMPO/E (HOMAIN);	Homo sapiens chromosome 21 segment HS21 0004	Homo saplens chromosome 21 segment HS2; C004	Homo sapiens KIAA0952 protein (KIAA0952), mRNA	Homo saplens KIAA0952 protein (KIAA0952), mRNA	Human beta-prime-adaptin (BAM22) gene, excn 16	Homo saplens E1A binding protein p300 (EP300) mRNA	Homo sapiens KIAA0952 protein (KIAA0952), ImRNA	601433525F1 NIH_MGC_72 Homo saplens cDNA clone IMAGE:3918607 5	601495208F1 NIH_MGC_70 Homo saplens cDNA clone IMAGE:3897457 5'	601496208F1 NIH_MGC_70 Homo sapiens cDNA clone IMAGE:3897457 5'	Homo sapiens mRNA for KIAA1363 protein, partial cds	Homo sapiens differentially expressed in FDCP (mouse homolog) 6 (DEF6), mRNA	Homo sapiens differentially expressed in FDCP (mouse homolog) 8 (DEF6), mRNA	oz09c07.x1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:1674828 3'	zv78a11.r1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA clone IMAGE:759740 5	zv78a11.r1 Soares_total_fetus_Nb2HF8_9w Homo sepiens cDNA clone IMAGE:769740 5'	602021846F1 NCL_CGAP_Brn67 Hamo sapiens cDNA clone IMAGE:4157339 5
Top Hit Database Source	EST_HUMAN	Ę	EST_HUMAN	EST_HUMAN	THUMAN		EST_HUMAN		EST_HUMAN	EST_HUMAN	EST_HUMAN		EST_HUMAN		EST_HUMAN		HOMAN	NT	NT			N			EST_HUMAN	EST_HUMAN	EST HUMAN	Z Z	뉟		EST_HUMAN	EST_HUMAN		EST_HUMAN
Top Hit Acession No.	BE274696.1		0.0E+00 AV738288.1	0.0E+00 AV738288.1		119828.1	3F344434.1	BE748899.1	BF377897.1	3F377897.1	3F313617.1		3E018750.1		0.0E+00 AA042813.1			0.0E+00 AL183204.2	0.0E+00 AL163204.2	7662401 NT	7662401 NT	J36264.1	4557556 NT	7662401 NT	3E895281.1	3E905563.1	0.0E+00 BE905563.1		11545748 NT	11545748 NT	A1076404.1	4A429001.1	4A429001.1	BF347039.1
Most Similar (Top) Hit BLAST E Value	0.0E+00 B	0.0E+00 D87685.1	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 B	0.0E+00	0.0E+00 B	0.0E+00 B		0.0E+00 B		0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00
Expression Signal	1.02	1.09	10.48	10.46	1.4	6.0	17.1	11.29	2.35	2.35	1.77		1.92		0.92		0.92	2.32	2.32	2.63	2.63	1.04	7.56	1.44	1.09	78.0	78.0	1.36	4.16	4.16	2.51	2.21	2.21	2.23
ORF SEQ ID NO:	21922	21925	21926	21927	21929			21934	21937				21948		21947			21956		21958			21984	21988	21996	21899	22000	22002	_	22035	22038		22039	
Exen SEQ (D NO:	12026	12028	12029	12029	12031	12033			12040	12040	12714		12046		12047		_	12055	12055			12061	12080	12086	1	12096	12096	12098				12140	12140	
Probe SEQ ID NO:	2138	2140	2141	2141	2143	2145	2148	2149	2152	2152	2156		2159		2160		2160	2168	2168	2169	2169	2174	2183	2199	2208	2209	2209	. 2212	2253	2253	2254	2258	2256	2258

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טווקום באטון רוטנסט באלים מספט ווווסמון	Top Hit Descriptor	Homo sapiens potassium channel Kv2.1 mRNĄ, complete cds	Homo sapiens flavin containing monooxygenas 3 (FMO3), mRNA	7f22s02.x1 NCI_CGAP_CLL1 Homo sepiens cDNA clone IMAGE:3296370 3' similar to TR:094939 094939 KIAA0857 PROTEIN;	Homo sapiens phosphorylase kinase alpha subunit (PHKA2) gene, exon 32	b57c08x1 NCI_CGAP_Ut2 Homo sapiens cDNA clone IMAGE:22831823'	Homo saplens sperm specific antigen 2 (SSFA2), mRNA	Homo saplens sperm specific antigen 2 (SSFA2), mRNA	Homo saplens titin (TTN) gene, alternative splice products, partial cds	Homo sapiens titin (TTN) gene, alternative splice products, partial cds	Homo sapiens signal regulatory protein, beta, 1 (SIRP-BETA-1) mRNA	AU131142 NT2RP3 Homo sapiens cDNA clorie NT2RP3002064 5'	601586843F1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:3941003 5'	MR1-SN0033-120400-002-a04 SN0033 Home sapiens cDNA	Homo sapiens KIAA0244 protein (KIAA0244), mRNA	Homo sapiens hexose-6-phosphate dehydrogenase (glucose 1-dehydrogenase) (H6PD), mRNA	Homo sapiens hexose-6-phosphate dehydrogenase (glucose 1-dehydrogenase) (H6PD), mRNA	Homo sapiens cytochrome P450 polypeptide 43 (CYP3A43) gene, partial cds; cytochrome P450 polypeptide 4 (CYP3A4) and cytochrome P450 polypeptide 7 (CYP3A4) genes complete cytochrome P450 polypeptide 2 (CYP3A4) genes complete cytochrome P450 polypeptide 2 (CYP3A47) genes complete cytochrome P450 polypeptide 2 (CYP3A47) genes complete cytochrome P450 polypeptide 2 (CYP3A47) genes complete cytochrome P450 polypeptide 3 (CYP3A47) genes complete cytochrome P450 polypeptide 3 (CYP3A47) genes complete cytochrome P450 polypeptide 3 (CYP3A47) genes complete cytochrome P450 polypeptide 3 (CYP3A47) genes complete cytochrome P450 polypeptide 3 (CYP3A47) genes complete cytochrome P450 polypeptide 3 (CYP3A47) genes complete cytochrome P450 polypeptide 3 (CYP3A47) genes complete cytochrome P450 polypeptide 3 (CYP3A47) genes complete cytochrome P450 polypeptide 3 (CYP3A47) genes complete cytochrome P450 polypeptide 3 (CYP3A47) genes complete cytochrome P450 polypeptide 3 (CYP3A47) genes complete cytochrome P450 polypeptide 3 (CYP3A47) genes cytochrome 3 (CYP3A47) genes cytochrome 3 (CYP3A47) genes cytochrome 3 (CYP3A47) genes	Polypeptide 6 (CYP3A6) gene, partial cds	AU118082 HEMBA1 Homo sapiens cDNA clone HEMBA1002839 5'	AU118082 HEMBA1 Homo sapiens cDNA clone HEMBA1002839 5'	AU118082 HEMBA1 Homo sapiens cDNA clone HEMBA1002839 5'	MR0-BN0070-090600-029-412 BN0070 Homy sapiens cDNA	AU119582 HEMBA1 Homo sapiens cDNA clone HEMBA1006155 5'	ox60b02.x1 Sceres_NhHMPu_S1 Homo sapiens cDNA clone IMAGE:1660683 3' similar to TR:008682 008662 230KDA PHOSPHATIDYLINOSITOL 4-KINASE.	xx15f07,x1 Soares NFL T GBC S1 Homo sapiens cDNA clone IMAGE:2813221 3' similar to TR:054924	064924 EXO84.;	601432608F1 NIH_MGC_72 Homo seplens cDNA clone IMAGE:3918168 5	AB005622 HeLa cDNA (T.Noma) Homo saplens cDNA similar to adenylate kinase isozyme 2	Homo sapiens glutamate receptor, tonotropic, N-methyl D-aspartate 24 (GRIN2A) mRNA	Homo sapiens gene for cholecystokinin type-A receptor, complete cds	Homo sapiens gene for chalecystakinin type-A receptor, complete ads
פון ביוסאם פון	Top Hit Database Source	NT	ᅜ	EST HUMAN	K	EST_HUMAN	ZI.	N F	NT	뉟	뉟	EST_HUMAN	EST_HUMAN	EST_HUMAN	F	ᅜ	Z		Þ	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	FST HUMAN		EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	NT	칟
	Top Hit Acession No.	L02840.1	6325466 NT	BE676095.1		AI625542.1	5803178 NT	5803178 NT	AF058332.1	AF058332.1	5174678 NT	AU131142.1	BE794026.1	AW867076.1	7662017	4758497 NT	4758497 NT		AF280107.1	AU118082.1	AU118082.1	AU118082.1		AU119582.1			AW303998.1	BE895605.1	AB005622.1	6006002 NT	D85606.1	D85606.1
	Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	_	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	00+400		0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00
	Expression Signal	1.19	2.03	-	10.08	2.72	1.78	1.76	4.28	4.26	2.88	1.75	6.71	86'0	1.97	1.44	4.		2.31	7.57	75.7	7.57	0.98	1.34	3.64		0.98	2.03	1.69	5.63	2.48	2.48
	ORF SEQ ID NO:			22054			22061	22062		22073	22079	22083			22085						22089			22138			22141				22162	
	Exon SEQ ID NO:	12147	12148	12155				12164	12173	12173	12182	12185	12186	L	12188	12189	12189		12190		12191		12208	12243	12248		12249	12251	12262	12265	12268	12268
	Probe SEQ ID NO:	2263	2264	227	2274	2276	2280	2280	2291	2291	2300	2304	2305	2306	2307	2308	2308		2309	2310	2310	2310	2327	2363	2388		2369	2371	2382	2386	2390	2390

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Top Hit Descriptor	Homo sapiens immunoglobulin-like transcript 1c variant 4 (ILT1c) gene, exon 6	602018058F1 NCI_CGAP_Bm67 Hamo sapiens cDNA clone IMAGE:4153670 5	Homo sapiens collagen, type XII, alpha 1 (COL12A1), mRNA	CMD-MT0033-150600-428-h11 MT0033 Homo sapiens cDNA	CM0-MT0033-150600-428-h11 MT0033 Homo sapiens cDNA	602184558T1 NIH_MGC_42 Homo sapiens cDNA clone IMAGE:4300383 3'	ha04h04.x1 NCI_CGAP_Kid12 Home sapiens cDNA clone IMAGE:2872759 3'	UI-HF-BP0p-als-c-07-0-UI.r1 NIH_MGC_51 Homo sapiens cDNA clone IMAGE:3072780 5'	Homo saplens protein kinase, AMP-activated, sipha 2 catalytic subunit (PRKAA2) mRNA	Homo sapiens protein kinase, AMP-activated, alpha 2 catalytic subunit (PRKAA2) mRNA	RC3-ST0197-300300-016-c04 ST0197 Homo saplens cDNA	601592530F1 NIH_MGC_7 Homo sapiens cDNA done IMAGE:3946518 5	Homo sapiens death receptor 6 (DR6), mRNA,	UI-H-BI4-aoz-b-08-0-UI.s1 NCI_CGAP_Sub8 Homo sapiens cDNA clone IMAGE:3086535 3'	Homo saplens mRNA for membrane transport protein (XK gene)	Homo sapians platelet-derived growth factor receptor-like (PDGFRL) mRNA	601503356F1 NIH_MGC_70 Homo sapiens clJNA clone IMAGE:3905148 5'	Homo saplens similar to rat integral membrane glycoprotein POM121 (POM121L1), mRNA	RC4-HT0276-160200-013-d05 HT0276 Home saplens cDNA	Homo sapiens hypothetical protein FLJ20366 (FLJ20366), mRNA	Human Sec62 (Sec62) mRNA, complete cds	601508211F1 NIH_MGC_71 Homo saplens clDNA clone IMAGE:3909866 5'	601489241F1 NIH_MGC_69 Homo saplens clDNA clone IMAGE:3891371 5'	601489241F1 NIH_MGC_69 Homo saplens clDNA clone IMAGE:3891371 5	Homo sapiens adiican mRNA, complete cds	601064738F1 NIH_MGC_10 Homo sapiens clONA clone IMAGE:3451161 5'	AU143277 Y79AA1 Homo sapiens cDNA clone Y79AA1001673 5'	AU143277 Y79AA1 Homo sapiens cDNA clone Y79AA1001673 5'	601105312F1 NIH_MGC_16 Homo sepiens aDNA alane IMAGE:2987955 5	601105312F1 NIH_MGC_15 Homo sapiens cDNA clone IMAGE:2987956 5'	7q27h12.71 NCI_CGAP_GC6 Homo saplens cDNA clone IMAGE: 3' similar to TR: 000246 000246 HYPOTHETICAL 9.3 KD PROTEIN :	Homo sepiens adlican mRNA, complete cds	601173631F1 NIH_MGC_17 Hamo sepiens cDNA clone IMAGE:3529159 5'
Top Hit Database Source	LN	EST_HUMAN	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	TN	NT	EST_HUMAN	EST_HUMAN	LN.	EST_HUMAN	NT	N	EST_HUMAN	NT	EST_HUMAN	NT	LN	EST_HUMAN	EST_HUMAN	EST_HUMAN	ΤN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	FST HIMAN	LZ	EST_HUMAN
Top Hit Acession No.	4F106275.1	3F345274.1	7776	0.0E+00 BE831003.1	0.0E+00 BE831003.1	0.0E+00 BF569144.1	0.0E+00 AW466922.1	0.0E+00 AW501010.1	5453965 NT	5453965 NT		0.0E+00 BE795542.1	57038	0.0E+00 BF509482.1	732684.2	5453871 NT	0.0E+00 BE910378.1	7657468 NT	3E150865.1	8923340 NT	193239.1	0.0E+00 BE886490.1	0.0E+00 BE875511.1	0.0E+00 BE875511.1	0.0E+00 AF245505.1	0.0E+00 BE536921.1	0.0E+00 AU143277.1	l	3E292896.1	0.0E+00 BE292896.1	0 0E+00 BE223041 1	AF245505.1	0.0E+00 BE296613.1
Most Similar (Top) Hit BLAST E Value	0.0E+00 AF	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.05+00	0.0E+00	0.0E+00	0.0E+00 Z32684.2	0.0E+00	0.0E+00	0.0E+00	0.0E+00 B	0.0E+00	0.0E+00 U93239.1	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.05+00.0	0.0E+00	0.0E+00
Expression	2.28	0.98	3.95	0.87	0.87	2.27	2.66	3.45	2.08	2.08	2.35	16.13	1.18	1.8	2.25	6.67	1.99	1.96	55.78	1.14	3.21	1.64	4.39	4.39	76.0	1.27	3.8	3.8	1.19	1.19	68.0	7.65	1.05
ORF SEQ ID NO:	22172	22178	22185	22188	22189			22202	22219	22220		22233	21671	22234	22236	 	22239	22240	22241	22242	22243	22249	22255	22256	22259	22267	22274	22275	22276	22277	87666	22280	22304
Exan SEQ ID NO:	12276	12279	12287	12291		12296	12305	12307	12321				11792	12340	12343	12345	12347	12348	12349	12350	12351	12357		12361		12379	12383	l	12384	12384	12385	12388	12414
Probe SEQ ID NO:	2398	2402	2410	2414	2414	2419	2428	2430	2444	2444	2457	2462	2463	2484	2467	2469	2471	2472	2473	2474	2475	2481	2486	2486	2488	2504	2509	2509	2510	2510	2511	2514	2540

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Single Exon Probes Expressed in Heart

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Tcp Hit Descriptor	Homo sapiens mRNA for KIAA1415 protein, parital cds	Homo sapiens mRNA for KiAA1415 protein, partial cds	UI-H-BW1-amp-f-12-0-UI.s1 NCI_CGAP_Sub7 Hamo saplens cDNA clone IMAGE:3070631 3'	802152653F1 NIH_MGC_81 Homo sapiens cDNA clone IMAGE:4293612 5	601279873F1 NIH_MGC_39 Homo sapiens cDNA clone IMAGE:3621786 5	Homo saplens mRNA for KIAA1321 protein, partial cds	h19b08.x1 NCI_CGAP_Brn25 Homo sapiens ¿DNA clone IMAGE:2168055 3' similar to gb:L20977 CALCIUM-TRANSPORTING ATPASE PLASMA MEMBRANE, BRAIN ISOFORM 2 (HUMAN);	Homo saplens TATA box binding protein (TBP) associated factor, RNA polymerase II, I, 28kD (TAF2I)	(IIIXIVA)	Thom o sapiens minny for NIAA 1450 protein, partiel cos	OU 350108F1 NIT MGC / Ham saplens cuna cione image: 3544304 5	601590108F1 NIH_MGC_7 Homo saplens cDNA clone IMAGE:3944304 5'	601143722F1 NIH_MGC_15 Homo sapiens cDNA clone IMAGE:3051389 5'	601584930F1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:3839222 5'	Homo sapiens IMP (inosine monophosphate) dehydrogenase 1 (IMPDH1) mRNA	Homo sapiens titin (TTN) mRNA	Homo saptens Bruton's tyrosine kinase (BTK), alpha-D-galactosidase A (GLA), L44-like ribosomal protein (L44L) and FTP3 (FTP3) genes, complete cds.	Homo sapiens guanylate cyclase-activating protein 2 (GUCA1B) gene, exon 1	Homo saplens mRNA for KIAA0536 protein, pertial cds	AU133385 NT2RP4 Homo saplens cDNA clone NT2RP4001964 5	Human bullous pemphigold antigen (BPAG1) mRNA, complete cds	AU130403 NT2RP3 Hamo sapiens cDNA clone NT2RP3000779 6'	AU130403 NT2RP3 Hamo sapians aDNA clone NT2RP3000779 5	RC1-OT0088-220300-011-d07 OT0086 Hama saplens cDNA	7h15h05.x1 NCI_CGAP_Co16 Hamo sapiens cDNA clone IMAGE:3316089 3'	601298714F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:3628923 5'	601278373F1 NIH_MGC_39 Homo sapiens cDNA clone IMAGE:3610287 5'	Homo sapiens hypothetical protein FLJ11052 (FLJ11052), mRNA	EST188414 HCC cell line (matastasis to liver in mouse) II Homo septens cDNA 5' end similar to ribosomal protein L29	601589625F1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:3943591 5'	Human beta-primo-adaptin (BAM22) gene, excin 5
Top Hit Database Source	NT	N N	EST_HUMAN	EST HUMAN	EST_HUMAN	N.T	EST_HUMAN	<u> </u>	Ž.	2	ESI_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	N	N N	FZ	·	N	EST_HUMAN	FN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	Z	EST HUMAN	EST HUMAN	NT
Top Hit Acession No.	0.0E+00 AB037836.1	AB037836.1	0.0E+00 BF513835.1	0.0E+00 BF672818.1	0.0E+00 BE616695.1	0.0E+00 AB037742.1	0.0E+00 AI571737.1	0.000	IN DETAKE	U.UE+UU ABUS/859.1	0.0E+00 BE/85445.1	0.0E+00 BE795445.1	0.0E+00 BE293328.1	0.0E+00 BE792472.1	4504686 NT	4507720 NT	178027.1	0.0E+00 AF173227.1	0.0E+00 AB011108.1	0.0E+00 AU133385.1	0.0E+00 M69225.1	1U130403.1	AU130403.1	0.0E+00 AW887015.1	3F000018.1	3E383165.1	0.0E+00 BE531263.1	8922843	0.0E+00 AA316723.1	3E794884.1	0.0E+00 U36253.1
Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00/	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	00.100	0.05+00	0.0E+00/	0.0=+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 U78027.1	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 AU	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00
Expression Signal	2,37	2.37	3.85	2.8	1.16	1.34	0.97		227	97.6	1.03	1.03	1.1	10.42	2.46	7.02	1.09	5.19	1.17	0.98	1.41	1.21	1.21	1.29	1	3.25	2.74	1.74	8.72	0.88	3.59
ORF SEQ ID NO:		22319		22326		22332	22333						22339			22365		22369	22370	22373	22374	22376	22377	22380	22385	22388		22413		22443	
Exan SEQ ID NO:	12654	12654	12426	12432	12434	12440	12441	3	12442	12444	12440	12445	12448	12456	12465	12727	12476	12477	12481	12484	12485	12487	12487	12490	12494	12495	12496	12623	12555	12656	12662
Probe SEQ ID NO:	2553	2553	2554	2560	2562	2569	2570	į	707	2)07	5/07	2574	2577	2585	2595	2601	2608	2609	2613	2616	2617	2619	2619	2622	2626	2827	2628	2658	2690	2691	2698

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Single Exon Probes Expressed in Heart

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Töp Hit Descriptor	Homo sapiens neuregulin 1 (NRG1), transcript variant SMDF, mRNA	Homo sapiens skeletal muscle LIM-protein 1 (FFIL1) gene, complete cds	601591991F1 NIH_MGC_7 Homo saplens cDNA done IMAGE:3945983 5'	602155923F1 NIH_MGC_83 Homo sapiens cDNA done IMAGE:4297132 5'	601335485F1 NIH_MGC_39 Homo sapiens cDNA done IMAGE:3689564 5'	AV721647 HTB Homo saplens cDNA clone HTBBYE09 5'	Homo sapiens spermatogenesis associated PD/1 (KIAA0757) mRNA	Homo sapiens spermatogenesis associated PD1 (KIAA0757) mRNA	Homo sapiens hypothetical protein FLJ20477 (FLJ20477), mRNA	Homo sapiens hypothetical protein FLJ20477 (FLJ20477), mRNA	Homo sapiens hypertension-related calcium-regulated gene mRNA, complete cds	AV651066 GLC Homo sapiens cDNA clone GLCCLD07 3'	CM1-TN0141-250900-439-b08 TN0141 Homo sapiens cDNA	CM1-TN0141-250900-439-b08 TN0141 Homo sapiens cDNA	Homo saplens cerebellar degeneration-related protein (34kD) (CDR1) mRNA	Homo sapiens cerebellar degeneration-related protein (34kD) (CDR1) mRNA	601580903F1 NIH_MGC_9 Homo saplens cDNA clone IMAGE:3929472 5'	RC4-HT0587-170300-012-d11 HT0587 Homo sapiens cDNA	Homo saplens chromosome 21 segment HS21C001	UI-H-BW1-amw-e-07-0-UI.s1 NCI_CGAP_Sub7 Homo saplens cDNA clone IMAGE:3071340 3'	Homo sapiens chondroitin sulfate proteoglycan 4 (melanoma-associated) (CSPG4), mRNA	602085579F1 NIH_MGC_83 Homo sapiens cDNA clone IMAGE:4249915 5'	Homo sapiens protein tyrosine phosphatase, receptor type, T (PTPRT), mRNA	AV725534 HTC Homo saplens cDNA clone HTCCCA03 5	AV725534 HTC Homo saplens cDNA clone HTCCCA03 5'	au55d04.y/ Schneider fetal brain 00004 Homo sapiens cDNA clone IMAGE:2518663 5' similar to	602071957F1 NCI CGAP Bin87 Homo septems cDNA clone IMAGE 4274679 5	80145081251 NIH MGC 65 Homo senjens citina citina (MAGE 3854642 5)	AU131494 NT2RP3 Homo sapiens oDNA clone NT2RP3002672 5	AU131494 NT2RP3 Homo saplens cDNA clone NT2RP3002672 5'	600944794F1 NIH_MGC_17 Homo sapiens c JNA clone IMAGE:2960808 5	600944794F1 NIH_MGC_17 Hamo saplens d)NA clone IMAGE:2960808 5'	glycoprotein D=Duffy group antigen [human, blood, Genomic DNA, 3089 nt]
Top Hit Database Source	LN	TN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT TA	NT	NT	IN	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	TN	.	EST_HUMAN	EST_HUMAN	IN	EST_HUMAN	IN	EST_HUMAN	NT	EST_HUMAN	EST_HUMAN	144441 151 150	EST HIMAN	EST HIMAN	EST HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	L
Top Hit Acesslon No.	7669517 NT	AF110763.1	BE796376.1	BF680632.1	BE563433.1	AV721647.1	5174486 NT	5174486 NT	8923441 NT	8923441 NT	AF290195.1	AV651066.1	BF377897.1	BF377897.1	4757963 NT	4757963 NT	BE747193.1	BE176836.1 .	AL163201.2	BF514110.1	4503098 NT	BF677694.1	7427522 NT	AV725534.1	AV725534.1	A1070463 4	AIO/9103.1	RE872768 1	AU131494.1	AU131494.1	BE300344.1	BE300344.1	S76830.1
Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	_	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	_	Ī	0.0E+00	0.0E+00	0.0E+00	0.0E+00				0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00	00.100					-		
Expression Signal	1.08	10.23	10.37	3.2	13.51	1.28	2.17	2.17	6.0	6.0	2.2	15.67	1.72	1.72	3.21	3.21	2.2	0.97	1.3	3.47	1.07	4.95	1.73	9.56	9.56	44 45	1 97	29	1.11	1.11	10.1	10.1	4.6
ORF SEQ ID NO:	22454	22455	22462	22463	22467		22470	22471	22472	22473	22474		22475	22478				22488		22498				22515	22516		22521			22524	22525		19966
SEQ ID NO:	12564	12565	12571	12572	12731		12577	12577	H	12578				12581						12604	12610			12623	12623	42635	1	1_		L	L		10151
Probe SEQ ID NO:	2700	2701	2708	2709	2712	2713	2715	2715	2716	2716	2717	2718	2719	2719	2723	2723	2727	2730	2741	2742	2748	2754	2758	2761	2761	27.62	278B	27.67	2769	2769	2770	2770	2775

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Top Hit Descriptor	Homo sapiens BTRCP2 mRNA for F-box and WD-repeats protein isoform C, complete cds	Homo sapiens ALR-like protein mRNA, partial cds	Homo sapiens ALR-like protein mRNA, partial cds	Homo sapiens cytochrome P450, subfamily I (dioxin-inducible), polypeptide 1 (glaucoma 3, primary infantile) (CYP1B1) mRNA	Homo sapiens cylochrome P450, subfamily I (dioxin-inducible), polypeptide 1 (glaucoma 3, primary infantile)	H. sepiens serine hydroxymethyltransferase pseudogene	Homo sapiens 5-aminolevulinate synthase 2 (ALAS2) gene, complete cds	Homo sapiens mRNA for KIAA1527 protein, partial cds	Homo sapiens partial rpl3 gene for ribosomal protein L3, U82 snoRNA, U83a snoRNA and U83b snoRNA oness	Homo sapiens chromosome 21 segment HS21 0001	Human AHNAK nucleoprotein mRNA, 5' end	PM0-HT0343-281299-003-e02 HT0343 Homo sapiens cDNA	PM0-HT0343-281289-003-e02 HT0343 Homo sapiens cDNA	H.sapiens Id3 gene for HLH type transcription factor	Homo sapiens chromosome 21 segment HS21,0068	Human transglutaminase mRNA, complete cds	Homo sapiens gammma-cytoplasmic actin (ACTGP3) pseudogene	Homo sapiens gammma-cytoplasmic actin (ACTGP3) pseudogene	Novel human mRNA from chromosome 1, which has similarities to BAT2 genes	H.sapiens mRNA for nuclear DNA helicase II	Homo sapiens protocedherin alpha C1 (PCDH-alpha-C1) mRNA, complete cds	Homo sapiens eukaryotic translation elongation factor 1 alpha 1 (EEF1A1) mRNA	Homo saplens eukaryotic translation elongation factor 1 alpha 1 (EEF1A1) mRNA	Homo sapiens serine/threonine kinase 9 (STK9) mRNA	DKFZp586G0621_r1 586 (syncarym: hute1) Homo sapiens cDNA clane DKFZp586G0621	Homo sapiens KIAA0054 gene product; Helicase (KIAA0054), mRNA	Homo sapiens KIAA0054 gene product; Helicase (KIAA0054), mRNA	Homo sapiens chondroitin sulfate proteoglycan 4 (melanoma-associated) (CSPG4), mRNA	QV2-BT0638-130400-138-h03 BT0636 Hamo sapiens cDNA	QV2-BT0636-130400-138-h03 BT0636 Hamo sapiens cDNA	Homo sapiens chromosome 21 segment HS21C006
Top Hit Database Source	NT TA	ZI.	NT	Ę	1	- LV	LV.	N	E	Į.	NT	EST_HUMAN	EST_HUMAN	N		IN	TN	TN	LN	TN	LN	IN	IN	LN	EST_HUMAN	N	NT	LN	EST_HUMAN	EST_HUMAN	NT
Top Hit Acession No.	AB033281:1	AF264750.1	AF264750.1	4503202 NT	114 0000000	45022 KB5980 1	AF068624.1	AB040860.1	4.1238852.1	AL163201.2	M80902.1	3E154504.1	BE154504.1	X73428.1	AL163268.2	M98478.1	D50657.1	D50657.1	AL096857.1	Y10658.1	AF152303.1	4503470	4503470 NT	4507280 NT	AL047599.1	7661883 NT	7681883 NT	4503098 NT	BE081896.1	BE081898.1	AL163206.2
Most Similar (Top) Hit BLAST E Value	0.0E+00		0.0E+00	0.0E+00	10.0	0.01	_	_			_	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00
Expression Signal	<u>g</u> .	1.37	1.37	3.58	0	8 6	121	1:1	1 07	2.24	5.94	1.58	1.58	1.38	2.84	2.58	43.48	. 43.46	1.34	4.62	98'0	25.08	25.08	2.42	1.35	1.26	1.25	3.42	4.95	4.95	1.64
ORF SEQ ID NO:		20479		20784	20700	22534				22538	22541	22543	22544			22547	22551	22552	22555			22558	22557	22568	22572	22573	22574		22576	22577	22588
Exan SEQ ID NO:	12638		10849	10941	l	12735	1	1	<u>l</u>	١_	L	12751		<u> </u>	12755			1		12788	12767			12779	12782	12783	12783	12784	12786	L	12794
Probe SEQ ID NO:	2778	2784	2784	2789	0300	2805	2806	2808	2814	2815	2819	2822	2822	2824	2826	2828	2833	2833	2837	2838	2839	2840	2840	2851	2854	2855	2855	2856	2858	2858	2866

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Probe SEQ ID NO: 2867 2867 2867 2867 2867 2867 2867 2867

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l able 4 Single Exon Probes Expressed in Heart	Tcp Hit Descriptor	qf43f09.x1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1752809 3'	Homo sapiens neuropilin 2 (NRP2) gene, complate cds, alternatively spliced	Homo sapiens neuropilin 2 (NRP2) gena, complate cds, alternatively spliced	Homo sapiens prospero-related homeobox 1 (PROX1) mRNA	Homo saplens mRNA for PKU-alpha, partial cds	Homo sapiens KIAA0737 gene product (KIAA0737), mRNA	Homo sapiens calcium channel, voltage-dependent, gamma subunit 3 (CACNG3), mRNA	Homo sapiens calcium channel, voltage-dependent, gamma subunit 3 (CACNG3), mRNA	Homo sapiens chromosome 21 segment HS210046	Human displacement protein (CCAAT) mRNA	Homo saplens semenogelin I (SEMG1) mRNA	Homo sapiens membrane-bound aminopeptidase P (XNPEP2) gene, complete cds	Homo sapiens heat shock 70kD protein 1 (HSPA1A), mRNA	Homo sapiens heat shock 70kD protein 1 (HSPA1A), mRNA	Isoform 2 of a novel human mRNA from chromosome 22	Homo sapiens putative transcription factor CR53 (CR53) mRNA, partial cds	Homo sapiens transcription factor IGHM enhancer 3, JM11 protein, JM4 protein, JM5 protein, T54 protein, JM10 protein, A4 differentiation-dependent protein, triple LIM domain protein 6, and synaptophysin genes.	complete cds; and L-type calcium channel a>	Homo saplens Interleukin 2 receptor, beta (IL2RB) mRNA	Human germline gene 16.1 for ig lambda L-chain C region (IgL-C16.1)	Homo sapiens F-box protein FBL5 (FBL5) mRNA, complete cds	Homo sapiens melanoma-associated antigen (MAGE-C1) gene, complete cds	Homo saplens SWI-SNF complex protein p270 mRNA, partiál cds	Homo sapiens NOD1 protein (NOD1) gene, exons 1, 2, and 3	Homo sapiens KIAA0469 gene product (KIAA0469), mRNA	Homo sapiens olfactory receptor-like protein (OLFR 42B) gene, OLFR 42B-9110 allele, partial cds	Homo sapiens potassium voltage-gated channel, Shab-related subfamily, member 1 (KCNB1) mRNA	Human ferritin heavy chain mRNA, complete cds	Homo saplens mRNA for KIAA0549 protein, pertial cds	Homo sapiens mRNA for KIAA0549 protein, pertial cds	ye32f03 s1 Stratagene lung (#937210) Homo sapiens cDNA clone IMAGE:119453 3' simitar to SP:S28539 S29539 BASIC PROTEIN, 23K - ;
jle Exon Prob	Top Hit Database Source	EST_HUMAN	Г	NT TN			NT	NT	NT	LN	NT		NT				NT										TN		NT	IN	NT	EST_HUMAN
Sing	Top Hit Acession No.	AI149880.1	AF281074.1	AF281074.1	4506118 NT	AB004884.1	7662273 NT	5729755 NT	5729755 NT	AL163246.2	M74099.1	4506882 NT	AF195953.1	5579469 NT	5579469 NT	AL359403.1	AF017433.1		AF196779.1	4504664 NT					AF149773.1	7662139 NT	AF042075.1	4826783 NT	120941.1	AB011121.1	AB011121.1	194870.1
	Most Similer (Top) Hit BLAST E Value	0.0E+00	0.0E+00				0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00		_					_	_	0.0E+00	0.0E+00	0.0E+00	_	0.0E+00	0.0E+00	0.0E+00
	Expression Signal	1.13	0.84	0.84	0.97	2.15	1.33	1.59	1.59	0.89	1.04	0.82	4.85	7.86	7.86	5.08	2.02		1.74	68.0	2.79	1.85	1.89	2.85	3.97	4.21	1.64	3.45	19.64	1.79	1.79	9.41
	ORF SEQ ID NO:		22677			22680			22691		22714			22730	22731		22735			22739							22792			22836		22845
	Exan SEQ ID NO:	12870	12879	12879	12880												12942			12947	12968	12970	12974	12995	12998	13001	13002	13028		13040	13040	13048
	Probe SEQ ID NO:	2943	2922	2952	2953	7987	5364	2965	2965	2890	2892	3001	3006	3008	3008	3011	3014		3017	3019	3038	3043	3047	3068	3069	3074	3075	3102	3111	3115	3115	3123

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	Top Hit Descriptor	601878507F1 NIH_MGC_55 Homo sapiens cDNA clone IMAGE:4107433 51	wu12h10.x1 NCI_CGAP_GC6 Homo sepiens cDNA clone IMAGE:2516803 3'	H.saplens mRNA for gamma-glutamytransferase	H. sapiens mRNA for gamma-glutamyltransferase	Homo sapiens neuradn III (NRXN3) mRNA	Homo saplens neurexin III (NRXN3) mRNA	Homo sapiens interleukin 1 receptor, type I (IL1R1) mRNA	Homo sapiens titin (TTN) mRNA	Homo sapiens titin (TTN) mRNA	Homo sapiens nucleolar phosphoprotein B23 (NPM1) mRNA, complete cds	Homo sapiens solute carrier family 25 (mitochondrial carrier; adenine nucleotide translocator), member 5 (SLC25A5), nuclear gene encoding mitochondrial protein, mRNA	Homo sapiens CREB binding protein (Rubinstein-Taybi syndrame) (GREBBP) mRNA	Homo sapiens CREB binding protein (Rubinstein-Taybi syndrame) (CREBBP) mRNA	ae97b11.s1 Stratagene schizo brain S11 Homo sapiens cDNA clone IMAGE:971133 3'	Homo saplens angiostatin binding protein 1 mRNA, complete cds	Homo sapiens angiostatin binding protein 1 mRNA, complete cds	Homo sapiens fibrillin 1 (Marfan syndrome) (FBN1) mRNA	Homo saplens titin (TTN) mRNA	Human connextn 43 processed pseudogene	Homo sapiens HLA class ill region containing jenascin X (tenascin-X) gene, partial ods; cytochrome P450 21- hydroxylase (CYP21B), complement component C4 (C4B) G11, halicase (SKI2W), RD, complement factor B	(Bf), and complement component C2 (C2) genes,>	Homo sapiens very large G-protein coupled receptor-1 (VLGR1) mRNA, complete cds	Homo saplens A kinase (PRKA) anchor protein 1 (AKAP1), mRNA	Homo sapiens A kinase (PRKA) anchor protein 1 (AKAP1), mRNA	Homo sapiens SWI-SNF complex protein p270 mRNA, partial cds	Homo sapiens hypothetical protein FLJ20695 (FLJ20695), mRNA	tr58f08.x2 NCI_CGAP_Pan1 Homo sapiens ciDNA clone IMAGE:2222535 3' similar to SW:RL11_RAT possage and properties to SW:RL11_RAT	Home sanians telomerase reverse transcriptase (TERT) gene exons 1-8	Home sanions telemenase reverse transcriptase (TERT) dene exons 1-8	Homo sapiens hormonally upregulated neu tumor-associated kinase (HUNK), mRNA	Homo sapiens hormonally upregulated neu tumor-associated kinase (HUNK), mRNA
	Top Hit Database Source	EST_HUMAN	EST_HUMAN	L	Ϋ́	LN	TN	N FA	ΝT	TN	TN	TN	LN LN	LN	EST_HUMAN	NT	LN	NT	NT	NT	ie Te	TN	NT	NT	TN	LΝ	NT	ECT UINANI	TN	12		ΙΝ
	Top Hit Acession No.	BF243336.1	Al968086.1	X98922.1	X98922.1	4758827 NT	4758827 NT	. 4504658 NT	4507720 NT	4507720 NT	M28699.1	4502098 NT	4758055 NT	4758055 NT	AA774783.1	AF286598.1	AF286598.1	4557590 NT	4507720 NT	M65189.1		AF019413.1	AF055084.1	4502014 NT	4502014 NT	AF265208.1	8923624 NT	A1690004.4		T	57213	7657213 NT
	Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	O.0E+00	_	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	_	_	0.0E+00	0.0E+00	0.0E+00	_	0.0E+00 /	0.0E+00	0.0E+00	0.0E+00		0.0E+00	700-20				0.0E+00
	Expression Signal	1,1	1.03	3.99	3.89	1.5	1.5	7.73	3.26	3.26	2.44	2.23	0.82	0.82	26.3	4.53	4.63	1.58	3.35	4.39		1.19	4.15	3.46	3.46	2	0.89	90 7	80.0	800	3 -	+
	ORF SEQ ID NO:	22862		22870		22883	22884	22892	22893	22894	22910	22912	22918	22919					22942		!	22949	22951	22959	22960		22974	coocc		23012		
	Exan SEQ ID NO:	13083	13065	13070		13081	13081	13088	13089	13089	13105	13108	13113	13113	13115	13123	13123		13139	13148			13151	15068	15068	13175	13176	40000	_	ł		13212
	Probe SEQ ID NO:	3138	3140	3145	3146	3156	3156	3163	3164	3164	3180	3183	3188	3188	3190	3198	3198	3210	3215	3224		3225	3227	3237	3237	3252	3253	Caca	3289	2000	3280	3290

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Homo sapiens zinc finger protein 45 (a Kruppel-associated box (KRAB) domain polypeptide) (ZNF45) mRNA x77c11.x1 Soares_NhHMPu_S1 Homo sapiens cDNA clone IMAGE:1662356 3' similar to WP:T19B4.4 te35g12x1 Soares_NhHMPu_S1 Homo sapiens cDNA clone IMAGE:2088742 3' similar to TR:000498 Homo sapiens potassium voltage-gated channel, Isk-related family, member 2 (KCNE2) mRNA ab51112.r1 Stratagene lung carcinoma 937219. Homo sapiens cDNA clone IMAGE:844367 57 ab51112.r1 Stratagene lung carcinoma 937219. Homo sapiens cDNA clone IMAGE:844367 57 ab51112.r1 Stratagene lung carcinoma 937219. Homo sapiens cDNA clone IMAGE:844367 57 Homo sapiens mRNA for G protein-coupled inward rectifier potassium channel, complete cds Homo sapiens cell-line KG1 transcriptional regulatory protein p54 mRNA, complete cds Homo sapiens cell-line KG1 transcriptional regulatory protein p54 mRNA, complete cds Homo saplens heparan sulfate (glucosamine) 3-O-sulfotransferase 1 (HS3ST1) mRNA hi84g01.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2979024 3' hi84g01.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2979024 3 Homo sapiens gamma-glutamylcysteine synthetase (GLCLC) gene, partial cds 601143863F1 NIH_MGC_15 Homo sapiens ciDNA clone IMAGE:3051373 57 601143863F1 NIH_MGC_15 Homo sapiens ciDNA clone IMAGE:3051373 57 602084583F1 NIH_MGC_83 Homo sapiens cDNA clone IMAGE:4248598 5' Homo sapiens glyceraldehyde-3-phosphate dehydrogenase (GAPD), mRNA Homo sapiens butyrophilin, subfamily 3, member A3 (BTN3A3), mRNA Homo saplens WAVE2 mRNA for WASP-family protein, complete cds Human endogenous retroviral DNA (4-1), complete retroviral segment Homo sapiens semenogelin II (SEMG2) mRNA Homo sapiens homologous yeast-44.2 protein mRNA, complete cds Top Hit Descriptor Homo sapiens retinoblastoma-binding protein 2 (RBBP2) mRNA Homo saplens midline 1 (Opitz/BBB syndrome) (MID1) mRNA Homo sapiens midline 1 (Opitz/BBB syndrome) (MID1) mRNA QV0-CT0225-230300-169-e01 CT0225 Homo sapiens cDNA 000498 MYASTHENIA GRAVIS AUTOANTIGEN GRAVIN Homo sapiens KIAA0806 gene product (KIAA0806), mRN Homo sapiens mRNA for KIAA1476 protein, partial cds Homo sapiens chromosome 21 unknown mRNA Novel human gene mapping to chomosome X CE13742; EST_HUMAN NT EST_HUMAN EST_HUMAN EST HUMAN EST HUMAN EST HUMAN EST_HUMAN EST_HUMAN HUMAN EST_HUMAN HUMAN Top Hit Database Source 4508028 NT 91.1 EST EST 뉟 눌 눌 4506884 NT 눋 z 눋 Þ z 6325463 NT 7682319 NT 4557752 NT 4826795 4826763 7689491 4557752 Top Hit Acession 4826967 AW852217.1 AW 664693.1 AW664693.1 BE304791.1 0.0E+00 AF078868.1 0.0E+00 AF045452.1 0.0E+00 AF231922.1 0.0E+00 AA626677.1 BE304791.1 0.0E+00 AF118846.1 0.0E+00 AI384007.1 AL 133204.1 0.0E+00|AB040909.1 0.0E+00 AI081907.1 AA626677. ġ BF676393. 0.0E+00 AF045452. AA626677 0.0E+00 (Top) Hit BLAST E Value 8 8 1.39 0.98 2.06 1.23 1.28 4.53 7.43 1.13 28.67 0.9 4 1:1 233 4. 0.96 98 0.95 0.98 0.79 0.79 Expression 23164 23172 23175 23176 23177 23182 23180 23183 23185 23194 23225 23242 23270 23274 23282 ORF SEQ 23191 23217 23226 23277 Ö N O SEQ ID 13365 13374 13386 13412 13485 13514 13357 13357 13414 13423 13445 13456 13480 13483 13492 13492 13371 13371 13377 13379 13457 1337 3455 3455 3455 3461 3448 Probe SEQ ID 3578 3600 3816 3440 3440 3473 3495 3497 3505 3529 3533 3540 3541 3564 3566 3569 3578 3596 3461 3463 3568 3571 ÿ

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NHTBCae15909f1 Normal Human Trabecular Bone Cells Homo sapiens cDNA clone NHTBCae15g09 NHTBCae15g09f1 Normal Human Trabecular Bone Cells Homo sapiens cDNA clone NHTBCae15g09 Homo sapiens ATP-sensitive Inwardly rectifying K-channel subunit (KCNJ6/BIR1) gene, complete cds Homo sapiens methyl CpG binding protein 2 (MECP2), mRNA UI-H-BW0-ajs-e-12-0-UI.s1 NCI_CGAP_Sub6 Homo sapiens cDNA clone IMAGE:2733022 3 UI-H-BW0-ajs-e-12-0-UI.s1 NCI_CGAP_Sub6 Homo sapiens cDNA clone IMAGE:2733022 3 Horno sapiens smooth muscle myosin heavy chain SM1 mRNA, alternatively spliced, partial Homo saplens similar to rat integral membrane glycoprotein POM121 (POM121L1), mRNA Homo sapiens similar to rat integral membrane glycoprotein POM121 (POM121L1), mRNA Homo sapiens similar to rat integral membrane glycoprotein POM121 (POM121L1), mRNA aa06g01.r1 Soares_NhHMPu_S1 Homo saplens cDNA clone IMAGE:812496 5' similar to SW.KRB4_SHEEP P02445 KERATIN, HIGH:SULFUR MATRIX PROTEIN, IIIB4. [1] Homo sapiens v-ets avian erythroblastosis virus E26 oncogene related (ERG), mRNA Homo sapiens v-ets avian erythroblastosis virus E26 oncogene related (ERG), mRNA Homo sapiens soluble neuropilin-1 mRNA, complete cds e6210.x1 Soares_NFL_T_GBC_S1 Homo sepiens cDNA clone IMAGE:2091307 3' Homo sapiens matrix metalloproteinase 24 (membrane-inserted) (MMP24), mRNA Homo sepiens protocadherin beta 3 (PCDH-beta3) mRNA, complete cds Homo sapiens DNA mismatch repair protein (MLH3) gene, complete cds Homo sapiens SH2-containing protein Nsp2 mRNA, complete cds Homo sapiens SH2-containing protein Nsp2 mRNA, complete cds Top Hit Descriptor Pan troglodytes offactory receptor (PTR208) gene, partial cds Homo sapiens KIAA0569 gene product (KIAA(569), mRNA Homo sapiens ribosomal protein S2 (RPS2) mRNA Homo saplens mRNA for KIAA0796 protein, partial cds Homo sapiens mRNA for KIAA1414 protein, partial cds Homo sapiens chromosome 21 segment HS21 0004 Homo saplans chromosome 21 segment HS21 C004 /us musculus junctophilin 1 (Jp1-pending), ஈRNA Human gene for Type XIX collagen at chain, exon Homo sapiens desmoplakin (DPI, DPII) (DSP) EST_HUMAN NT **EST HUMAN** EST_HUMAN EST_HUMAN HUMAN EST_HUMAN Top Hit Database Source EST 눋 눋 눈 눋 7710148 NT 눋 ż 눋 z 4758189 NT 5729928 NT 4506718 NT 7657065 NT 7662183 7657065 7657468 10181139 7657468 7657468 Fop Hit Acession 0.0E+00 AW 298134.1 AW298134.1 0.0E+00 AA852743.1 0.0E+00 AA852743.1 AA463659.1 AF195658.1 0.0E+00 AF179733.1 AF152496.1 0.0E+00 AF124250.1 0.0E+00 AF124250.1 0.0E+00 AL163204.2 0.0E+00 AL163204.2 0.0E+00 AB004630.1 AB037835.1 4F145712.1 AF020091.1 0.0E+00 AI377699.1 AB018339. ģ 0.0E+00 S78685.1 0.0E+00/ 0.0E+00 / 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 (Top) Hit BLAST E Value Aost Similar 3.88 11.67 1.5 1.78 3.53 0.99 .03 3.35 0.91 7.88 1.02 1.02 1.13 1.01 2.3 69 1.69 1.45 4.46 3.26 222 Expression Signal 23446 23458 23453 23459 23318 23333 23344 23370 23402 23442 23447 23448 23455 23456 ORF SEQ ID NO: 23324 23331 23369 23401 23323 23377 23327 13548 13559 13618 13618 13673 13676 13533 13559 13613 13864 13537 13537 13540 13544 13582 13677 13671 SEQ ID 13591 1361 ÿ 3645 3699 3701 3705 3746 3747 3748 3751 3751 3752 3758 3758 3759 3763 3764 SEQ ID 3619 3619 3623 3628 3630 3632 3760 3668 3669 3623 3626 3677 ÿ

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Single Exon Probes Expressed in Heart

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	Top Hit Descriptor	Homo sapiens KIAA0569 gene product (KIAA0569), mRNA	Homo sapiens myosin light chain kinase Isoform 2 (MLCK) mRNA, complete cds	Homo sapiens myosin light chain kinase isoform 2 (MLCK) mRNA, complete cds	Homo sapiens 5-hydroxydryptamine (serotonin) receptor 1D (HTR1D) mRNA	Homo sapiens chromosome 21 segment HS21C079	Homo sapiens transient receptor potential channel 5 (TRPC5), mRNA	Homo sapiens chromosome X open reading frame 5 (CXORF5) mRNA	Homo sapiens chromosome X open reading frame 5 (CXORF5) mRNA	Human zinc finger protein ZNF134 mRNA, complete cds	Homo saplens potassium voltage-gated channal. Shab-related subfamily, member 1 (KCNB1) mRNA	Homo saplens familial mental retardation protein 2 (FMR2) gene, exon 11	Homo saplens SC35-interacting protein 1 (SRRP129), mRNA	Homo sapiens amphiphysin gene, partial cds	wk01f01.x1 NCI_CGAP_Lym12 Homo sapiens cDNA clone IMAGE:2411065 3' similar to TR:O43340 043340 R28830 - 'contains element PTR7 remetitive element -	Homo saplens ribosomal protein S8 (RPS8), mRNA	DKFZp434N0413_r1 434 (synonym: htes3) Hcmo saplens cDNA clone DKFZp434N0413 5	Homo sapiens AP1 gamma subunit binding protein 1 (AP1GBP1), mRNA	Homo sapiens AP1 gamma subunit binding protein 1 (AP1GBP1), mRNA	Homo sapiens glutamate receptor, metabotropic 3 (GRM3) mRNA	Homo sapiens melanoma antigen, family B, 1 (MAGEB1) mRNA	Homo sapiens HBP17 heparin-binding and FGF-binding protein gene, complete cds	Homo sapiens ryanodine receptor 3 (RYR3) mRNA	Homo sapiens zinc finger protein (KIAA0412) mRNA	RC3-HT0860-170800-011-a12 HT0860 Homo saplens cDNA	MXRA5 Human matrk tissue expression library Homo sapiens cDNA clone Incyte 1998726 similar to MXRA5 Matrk remodeling associated gene 5	sion library Homo sapiens cDNA clone Incyte 1996728 similar to MXRA5	Matrix remodeling associated gene 5	Homo sapiens F-box protein Fbl3b (FBL3B) mRNA, partial cds	601236966F1 NIH_MGC_44 Homo septens cDNA clone IMAGE:3606800 5	601153727F1 NIH_MGC_19 Homo saplens cDNA clone IMAGE:3509743.5'	PM3-LT0031-100100-003-h09 LT0031 Homo gaplens cDNA
	Top Hit Database Source	Ε	N	Ν	٦	N	۲	L	님	NT	LN	IN	FZ	LN	FST HIMAN	LN	EST_HUMAN	LN	TN	TN	TN	TN	IN	TN	EST_HUMAN	EST_HUMAN		EST_HUMAN	TN	EST_HUMAN	EST_HUMAN	EST_HUMAN
	Top Hit Acession No.	7662183 NT	AF069601.2	AF069601.2	4504534 NT	AL163279.2	6912735 NT	4503178 NT	4503178 NT	J09412.1	4826783 NT	AF012615.1	4759171 NT	4F099117.1	A1864727.1	4506742 NT	AL040338.1	6005887 NT	F005887 NT	4504138 NT	4505078 NT	AF149412.1	4508758 NT	4585642 NT	3F366285.1	0.0E+00 AW888221.1		0.0E+00 AW888221.1	1F129533.1	3E378602.1	0.0E+00 BE313146.1	0.0E+00 AW580740.1
	Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.05+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00 B	0.0E+00	0.0E+00/
	Expression Signal	2.39	1.31	1.31	1.97	1.61	1.12	5.67	5.67	3.93	1.63	66'0	1.87	0.82	2.54	8.08	1.41	+	-	2.45	1.8	0.87	1.27	1.62	1.75	1.4		1.4	1.78	3.2	1.27	0.97
	ORF SEQ ID NO:		23462	23463			23470		23479				23487	23489	23498	23502	23508	23515	23516	23518		23521	23534	23537	23544	23546	-	23547	23552	23557	23558	23563
	Excan SEQ ID NO:				13682				li	13694	13698	13699	13700	13702	13711	13714	13719	13725		1			13742	13745	13751	13753		13753	13759	13764	13765	13771
	Probe SEQ ID NO:	3765	3768	3768	3769	3773	3775	3780	3780	3782	3784	3787	3788	3780	3799	3802	3807	3813	3813	3815	3816	3820	3830	3833	3840	3842		3842	3848	3853	3854	3860

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Top Hit Descriptor		Homo sapiens cancer-testis antigen CT10 (CIT10) gene, complete cds	Homo sapiens cancer-testis antigen CT10 (CIT10) gene, complete cds	Human MHC class II lymphocyte antigen DPw4-beta-2 pseudogene, exon 2	Homo sapiens chromosome 21 segment HS21C103	Novel human gene mapping to chomosome 20	Homo sapiens chromosome 21 segment HS21C084	Homo sapiens chromosome 21 segment HS21C068	Homo saplens eukaryotic translation elongation factor 1 alpha 1 (EEF1A1) mRNA	Homo sapiens KIAA0569 gene product (KIAA0569), mRNA	Human zinc finger protein ZNF133	Chlorocebus aethiops mRNA for ribosomal protein S4X, complete cds	Homo sapiens mRNA for UGA suppressor tRNA-associated antigenic protein (tRNA48 gene)	Homo sapiens mRNA for rapa-2 (rapa gene)	Homo saplens mRNA for rapa-2 (rapa gene)	Homo sapiens retinoblastoma-binding protein 4 (RBBP4) mRNA	Homo sapiens retinoblastoma-binding protein 4 (RBBP4) mRNA	Homo sapiens phosphoribosylgiycinamide formytransferase, phosphoribosylgiycinamide synthetase, phosphoribosylamindarole synthetase (GART) mRNA	Homo sapiens G protein-coupled receptor 21 (GPR21), mRNA	Homo sapiens mRNA for KIAA0287 gene, partial cds	Homo sapiens ras GTPase activating protein-like (NGAP) mRNA	Homo sapiens IMP (Inosine monophosphate) dehydrogenase 1 (IMPDH1), mRNA	Novel human mRNA from chromosome 1, which has similarities to BAT2 genes	Homo sapiens DGCR8 (DGCR8) mRNA, complete cds	Homo sapiens DNA polymerase zeta catalytic subunit (REV3) mRNA, complete cds	Homo saplens protein kinase, X-linked (PRKX) mRNA	Homo sapiens protein kinase, X-linked (PRKX) mRNA	Homo sapiens butyrophilin, subfamily 3, member A2 (BTN3A2), mRNA	Homo sapiens GA-binding protein transcription factor, alpha subunit (60kD) (GABPA), mRNA	Homo sapiens GA-binding protein transcription factor, alpha subunit (60kD) (GABPA), mRNA	Homo sapiens hypothetical protein FLJ10379 (FLJ10379), mRNA	Homo sapiens hypothetical protein FLJ10379 (FLJ10379), mRNA	wu04d04.x1 NCI_CGAP_GC8 Homo sapiens cDNA clone IMAGE:25159753'	wu04d04.x1 NCI_CGAP_GC8 Homo saplens cDNA clone IMAGE:25159753'
	Source	NT	NT	N	NT	N	Z	뒫	NT	Ę	M	١	١	F	N	ΙN	NT	TN	L _N	뉟	N	IN	L	LN	LN	LN	TN	LN	NT	NT	LN	L	EST_HUMAN	EST_HUMAN
Top Hit Acession	•	AF116195.1	AF116195.1	M23910.1	0.0E+00 AL163303.2	0.0E+00 AL118494.1	0.0E+00 AL163284.2	0.0E+00 AL163268.2	4503470 NT	7662183 NT	U09366.1	0.0E+00 AB015610.1	0.0E+00 AJ238617.1	AJ277276.1	0.0E+00 AJ277278.1	5032028 NT	5032026 NT	4503914 NT	4885306 NT	B00662	4758807 NT	11419297 NT	AL096857.1	0.0E+00 AF165527.1	AF157476.1	4828947 NT	4826947 NT	5901905 NT	4503854 NT	4503854 NT	8922391 NT	8922391 NT	A1982597.1	A1982597.1
Most Similar (Top) Hit	Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 A	0.0E+00 AI
Expression		4.6	4.6	3.83	5.44	1.53	2.89	1.46	25.75	1.18	2.05	6.24	3.79	3.28	3.28	5.63	6.63	0.81	4.89	1.94	1.28	6.1	1.58	2.7	0.85	0.8	0.8	66.0	1.16	1.16	1.05	1.05	4.59	4.59
ORF SEQ		23587					23605	23613				23845						23677		23683		23685			23697	20872	20873			i			23722	23723
S	Ö.	13803			13815	13822	13825	13833		1		13867	H				13888	13900		13907					- [- 1		ŀ	13944	13944
Probe SEQ ID	Ö	3893	3893	3903	3905	3912	3916	3924	3935	3939	3940	3960	3968	3976	3976	3981	3981	3993	4000	4001	4003	4004	4005	4013	4017	4022	4022	4028	4029	4029	4032	4032	\$	4041

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Homo sapiens transglutaminase 3 (E polypeptide, protein-glutamine-gamma-glutamyfiransferase) (TGM3) qd23106.x1 Soares_placenta_8to9weeks_2NbHP8to9W Homo sapiens cDNA clone IMAGE:1724579 3' ba51f04.x1 NIH_MGC_10 Home sapiens cDNA clone IMAGE:2900095 3' similar to SW:THI2_BOVIN Q95108 MITOCHONDRIAL THIOREDOXIN PRECURSOR; Homo sapiens polycystic kidney disease (polycystin) and REJ (sperm receptor for egg jelly, sea urchin zu68h07.s1 Soares_testis_NHT Homo sapier;s cDNA clone IMAGE:743197 3' similar to contains Alu zu88h07.s1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:743197 3' similar to contains Alu JI-HF-BM0-edx-c-02-0-UI.r1 NIH_MGC_38 Hamo saplens cDNA clone IMAGE:3063147 5 601120778F1 NIH_MGC_20 Homo sapiens cDNA clone IMAGE:2967690 5 Homo sapiens proprotein convertase subtilisin/kaxin type 2 (PCSK2) mRNA Homo sapiens hypothetical protein FLJ10498 (FLJ10498), mRNA lomo sapiens hypothetical protein FLJ10498 (FLJ10498), mRNA domo sapiens mRNA for olfactory receptor protein, pseudogene **Top Hit Descriptor** fomo sapiens nuclear receptor coactivator 3 (NCOA3), mRNA repetitive element; contains element MER35 repetitive element; MR1-HT0707-100500-001-a02 HT0707 Homo sapiens cDNA MR1-HT0707-100500-001-a02 HT0707 Homo saplens cDNA repetitive element; contains element MER35 repetitive element; Human apolipoprotein B-100 mRNA, complete cds PM2-DT0023-080300-004-a08 DT0023 Homo sapiens cDNA Homo sapiens myelodysplasia syndrome 1 (MiDS1) mRNA Homo sapiens F-box protein Fbl4 (FBL4) mRNA, partial ods Homo sapiens myelodysplasia syndrome 1 (MIDS1) mRNA similar to contains MER20.b2 MER20 repetitive element; Homo sapiens desmoplakin (DPI, DPII) (DSP) mRNA Homo sapiens desmoplakin (DPI, DPII) (DSP) mRNA Homo sapiens protein kinase C, nu (PRKCN), mRNA Homo sapiens chromosome 21 segment HS21C103 Homo saplens semenogelin I (SEMG1) mRNA Human CBFA3 (Cbfa3) gene, partial cds Homo sapiens titin (TTN) mRNA homolog)-like (PKDREJ) mRNA Homo sapiens titin (TTN) mRNA HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST HUMAN EST_HUMAN Top Hit Database Source EST_HUMAN EST HUMAN HUMAN EST EST 4507476 NT 5174632|NT 눋 8922466 NT 4758199 NT 4758199 NT 눋 눌 4826827 NT 4505646|NT 4506882 NT 4826827 NT 6563384 NT 4507720 4507720 8922468 Top Hit Acession AW408788.1 0.0E+00 BE184856.1 0.0E+00 BE184856.1 0.0E+00 BE274217.1 0.0E+00 AW675599.1 0.0E+00 AW408788.1 0.0E+00 AW93689.1 AL163303.2 0.0E+00 AA401438.1 AA401438.1 0.0E+00 AI189844.1 AF174590.1 AJ003145.1 ġ 0.0E+00 J02610.1 0.0E+00|A 0.0E+00 A 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 (Top) Hit BLAST E Value 88.0 6.19 0.88 0.88 0.98 0.88 0.97 86: .52 1.33 86. 7.82 0.87 8.21 0.88 7.84 0.95 1.92 0.87 1.99 .33 8.21 Expression Signal 23733 23837 23851 23868 23876 23725 23726 23751 23775 23792 23801 23803 23804 23897 23904 23877 23752 ORF SEQ 23734 ΘNÖ 13946 14105 SEQ ID 13946 13957 13966 13973 14122 14128 13958 13983 13998 14013 14026 14030 14063 14076 14090 14098 14098 13950 13971 13973 14013 14030 14037 14096 ö 4210 4044 4055 4176 Probe SEQ ID 400 400 400 400 400 4113 4113 4128 4130 4137 4163 4190 4188 4186 4198 4224 4071 4071 4098 4098 4081 ğ

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	Top Hit Descriptor	Homo saplens protein kinase C, nu (PRKCN), mRNA	Human G2 protein mRNA, partial cds	Human G2 protein mRNA, partial cds	Homo saplens COMPLEMENT COMPONENT C1q RECEPTOR (C1QR), mRNA	Homo saplens gap junction protein connextn-36 (CX38) gene, complete cds	Human Ig light chain VL1 region germline (humlv1c2c) gene, partial cds	Homo sapiens plasma membrane calcium ATPase isoform 1 (ATP2B1) gene, alternative splice products,	peruei cos	H.sapiens H2B/h gene	H.sapiens H2B/h gene	H.sapiens H4/d gene for H4 histone	H.sapiens H4d gene for H4 histona	Homo sapiens KIAA0390 gene product (KIAA0390), mRNA	Homo sapiens KIAA0390 gene product (KIAA0390), mRNA	Homo sapiens caudal type homeo box transcription factor 4 (CDX4), mRNA	Homo saplens mRNA for KIAA1360 protein, partial cds	Homo sapiens myosin regulatory light chain interacting protein (MIR), mRNA	Homo saplens membrane-bound aminopeptidase P (XNPEP2) gene, complete cds	Homo saplens ACTN2 gene for alpha-Actinin/2, exon 10	Homo saplens ACTN2 gene for alpha-Actinin 2, exon 10	Homo saplens HPS1 gene, intron 5	xx68e08.x1 NCI_CGAP_Eso2 Homo sapiens cDNA clone IMAGE:2589446 3' similar to SW:AHNK_HUMAN Q09666 NEUROBLAST DIFFERENTIATION ASSOCIATED PROTEIN AHNAK;	Homo sapiens LIM domain kinase 2 (LIMK2), transcript variant 2a, mRNA	Homo sapiens vascular endothelial cell growth factor 165 receptor/neuropilin (VEGF185) mRNA, complete	ods	Hamo sapiens chromosome 21 segment HS21C007	PM1-HT0305-101199-002-403 HT0305 Homo sapiens cDNA	Homo sapiens mRNA for putative ankyrin-repeat containing protein (ORF1)	Homo sapiens mRNA for putative ankyrin-repeat containing protein (ORF1)	Homo saplens G protein-coupled receptor 50 (GPR50) mRNA	Homo saplens serine-threonine protein kinase (MNBH) mRNA, complete cds	H.saplens pancreatic polypeptide receptor PPI gene	Homo sapiens ATP-sensitive inwardly rectifying K-channel subunit (KCNJ8/BIR1) gene, exon
	Top Hit Database Source	ΙN	NT	IN	Ę	N L	INT	1	Z	LN_	LN	LΝ	NT	NT	F	ΙΝ	NT	F	ΤN	NT	NT	N	EST HUMAN	FZ		LN	LN	EST_HUMAN	TN	IN.	TN	NT	NT	NT
	Top Hit Acession No.	6583384 NT	J10991.1	U10991.1	6912281 NT	AF153047.2	U03901.1	, ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	L14561.1	Z80780.1	Z80780.1	X60483.1	X60483.1	7662091 NT	7662091 NT	4885126 NT	AB037781.1	7019456 NT	AF195953.1	AJ249765.1	AJ249765.1	AF200629.1	AW084964.1	8051619 NT		AF016050.1	AL163207.2	AW381570.1	AJ278120.1	AJ278120.1	4758467 NT	AF108830.1	Z66526.1	S78684.1
	Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00				_	0.0E+00	0.0E+00	0.0E+00	00+30.0	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00			_	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00
	Expression Signal	0.87	1.17	1.17	10.2	1.15	1.59		5.17	2.78	2.78	1.58	1.58	8.95	8.95	10.81	96.0	1.1	6.85	8.1	8.1	1.92	40.23	1.91		1.38	7.5	1.29	1.83	1.83	3.91	2.3	1.47	1.12
	ORF SEQ ID NO:	23905	23911		23917		23943			23955	23956	23962		23969		23982		24012		24025			24080			24083			24094	24095	24097	24098		24111
	Exon SEQ ID NO:	14128	14135	14135	14144	14162	14167		1	14177	14177	14183	14183	14187	14187	14197	14200	14230	14238	14243	14243	14264	14296	15073	i	- 1	14302	14304	14310	14310	14312	14313	L_{-I}	14324
	Probe SEQ (D NO:	4230	4237	4237	4245	4263	4268		4274	4278	4278	4284	4284	4289	4289	4289	4302	4333	4341	4346	4346	4368	4401	4403		4405	4408	4410	4416	4416	4418	4419	4424	4429

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	Top Hit Descriptor	QV2-BT0635-160400-142-h05 BT0635 Hamo sapiens cDNA	Homo saplens truncated tenascin XB (TNXB) gene, partial cds and TNXA gene recombination breakpoint region	Homo saplens mRNA for KIAA1399 protein, partial cds	Homo saplens mRNA for KIAA1399 protein, partial cds	Human displacement protein (CCAAT) mRNA	Homo saplens butyrophilin, subfamily 2, member A2 (BTN2A2), mRNA	Homo sapiens butyrophilin, subfamily 2, member A2 (BTN2A2), mRNA	ya83g04.r2 Stratagene fetal spleen (#937205) Homo sapiens cDNA clone IMAGE 88310 5	ya83g04.r2 Stratagene fetal spleen (#937205) Homo sapiens cDNA clone IMAGE:68310 5	601158935F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3505521 6'	601285246F1 NIH_MGC_44 Homo sapiens cONA clone IMAGE:3607067 5'	Human AHINAK nucleoprotein mRNA, 5' end	Human haptoglobin and haptoglobin-related protein (HP and HPR) genes, complete cds	Human haptoglobin and haptoglobin-related protein (HP and HPR) genes, complete cds	Homo saplens cyclophilin-related protein (NKTR) gene, complete cds	Homo sapiens KiAA0563 gene product (KiAA0563), mRNA	Human CYP2D7AP pseudogene for cytochroine P450 2D8	Homo sapiens bromodomain adjacent to zinc finger domain, 2B (BAZ2B), mRNA	Homo saplens bromodomain adjacent to zinc (inger domain, 2B (BAZ2B), mRNA	Homo sapiens alpha-3 type IX collagen (COL9A3) gene, promoter region, and exons 1-28	Homo sapiens proteinx0008 (AD013), mRNA	Homo saplens proteinx0008 (AD013), mRNA	UI-H-BI3-alw-c-04-0-UI.s1 NCI_CGAP_Sub5 Homo sapiens cDNA clone IMAGE:2733294 3'	Homo sapiens aldehyde dehydrogenase 12 (ALDH12) mRNA, complete cds	Homo saplens HSPC024-iso mRNA, complete cds	Human connexin 43 processed pseudogene	Homo saplens glutathione S-transferase theta 2 (GSTT2) and glutathione S-transferase theta 1 (GSTT1) genes. complete cds	M.fascicularis mRNA for metalloprotease-like, disintegrin-like protein. IVa	Homo sapiens Williams-Beuren syndrome deletion transcript 9 (WBSCR9) mRNA complete cds	Mus musculus zinc finger transcription factor Kalso mRNA, complete cds	Homo sapiens fregile X mental retardation 2 (FMR2) mRNA	Homo sapiens actin, alpha, cardiac muscle (ACTC), mRNA
-	Top Hit Database Source	EST HUMAN	LN TN	NT	NT	NT	NT	NT	EST HUMAN	EST HUMAN	EST_HUMAN	T_HUMAN	Г	L'N	LZI	N-I	N.	Ł	FZ		NT.			T_HUMAN		LN	L	L		LN	Z		
	Top Hit Acession No.	0.0E+00 BE081527.1	0.0E+00 AF086641.1	0.0E+00 AB037820.1		74099.1	6453812 NT	6453812 NT	T56945.1	T56945.1						2	7662181 NT		7304922 NT	7304922 NT	-026801.1	7019320 NT	7019320 NT	1	-303134.1	1.1		0.0E+00 AF24078B.1		-084479.1	0.0E+00 AF097416.1	4503766 NT	4885048 NT
	Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 M	0.0E+00	0.0E+00	0.0E+00	0.0E+00 T	0.0E+00	0.0E+00	0.0E+00 M80902.1	0.0E+00 M69197.1	0.0E+00 M69197.1	0.0E+00 AI	0.0E+00	0.0E+00 X58467.1	0.0E+00	0.0E+00	0.0E+00 A	0.0E+00	0.0E+00	0.0E+00/	0.0E+00	0.0E+00 AF	0.0E+00 M65189.1	0.0E+00	0.0E+00 X87205.1	0.0E+00 AF	0.0E+00	0.0E+00	0.0E+00
	Expression Signal	2:32	2.01	2.65	2.65	2.55	1.82	1.82	1.03	1.03	66.0	1.11	37.36	3.14	3.14	1.12	1.34	1.54	0.95	0.85	1.09	0.92	0.92	1.78	1.11	1.8	2.04	2.83	2.79	1.3	1.47	3.51	61.82
	ORF SEQ ID NO:	24246		24257	24258	24259	24262	24263	62661	19930		24267		24293		24297	24298			24320	24327	24330	24331	24355	24363		_		24408	24410	24411	24412	24414
	Exan SEQ ID NO:	14458	14465	14470	14470	14471	14474	14474	10108	10108	14475	14481	14502	14505	14505	14508	14509	14524	14532	14532	14538	14541	14541	14562	14587	14572	14583	14620	14622	14624	14825	14828	14828
	Probe SEQ ID NO:	4566	4574	4680	4580	4581	4585	4585	4586	4586	4587	4593	4614	4617	4617	4620	4621	4636	4644	4644	4652	4655	4655	4676	4681	4686	4697	4735	4737	4739	4740	4741	4743

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Top Hit Descriptor	ZINO FINGER PROTEIN 132	Homo sapiens hypothetical protein DKFZp762E1312 (DKFZp762E1312), mRNA	Homo sapiens chromosome 21 segment HS21C003	Homo sepiens hypothetical protein FLJ20073(FLJ20073), mRNA	Homo sapiens KIAA0187 gene product (KIAA0187), mRNA	Human Tor-C-delta gene, exons 1-4; Tor-V-dialta gene, exons 1-2; T-cell receptor alpha (Tor-alpha) gene, J1- J81 segments: and Tor-C-alpha gene, exons 1-4	Human Tor-C-delta gene, exons 1-4; Tor-V-delta gene, exons 1-2; T-cell receptor alpha (Tor-alpha) gene, J1-	H. sapiens MeCP-2 dene	H.sapiens MeCP-2 gene	Human collagenase type IV (CLG4) gene, excn 2	Hamo sapiens chromosome 21 segment HS21C080	Homo sapiens TATA box binding protein (TBP)-associated factor, RNA polymerase II, I, 28kD (TAF2I) mRNA	Homo sepiens low density lipoprotein-related protein 2 (LRP2), mRNA	H.sapiens MICA gene	Homo sapiens zinc finger protein (KIAA0412) mRNA	Homo sapiens mRNA for KIAA0633 protein, partial cds	Mus musculus zinc finger protein interacting with K protein 1 (Zik1), mRNA	Hamo sapiens meningioma expressed antigen 6 (colled-coll proline-rich) (MGEA6), mRNA	Hamo sapiens desmoplakin (DPI, DPII) (DSP) mRNA	Hamo sapiens gene encoding filensin, exan 8	Hamo sapiens zinc-finger DNA-binding protein (HUMHOXY1), mRNA	Homo sapiens mRNA for immunoglobulin kappa light chain, anti-RhD, therad 7	Homo sapiens MHC class 1 region	Homo sapiens opioid receptor, delta 1 (OPRD 1) mRNA	Homo saplens splice variant AKAP350 mRNA, partial cds	Homo sapiens titin (TTN) mRNA	Homo sapiens titin (TTN) mRNA	Homo saplens partial TTN gene for titin	Homo sapiens titin (TTN) mRNA	Homo sapiens COL4A6 gene for a6(IV) collagien, exon 44 and partial ods
Top Hit Database Source	SWISSPROT	N L	N	N	NT	LZ	TIV	L	Į.	NT	LN L	FZ	LN	N.	TN	FZ	LN	NT	LN	TN	LN	LN	NT	١N	NT	LN	LN	TN	TN	Į.
Top Hit Acession No.	P52740	8922180 NT	AL163203.2	R923080 NT	7661979 NT	V94081.1	40000	X94628.1	94628.1		0.0E+00 AL163280.2	5032150/NT	G806918 NT	K92841.1	4585642 NT	0.0E+00 AB014533.1	6677648 NT	5174560 NT	4758199 NT	/16723.1	7705546 NT	4J010442.1	0.0E+00 AF055066.1	4505508 NT	4F091711.1	4507720 NT	4507720 NT	0.0E+00 AJ277892.1	4507720 NT	
Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00 A	0.0E+00	0.0E+00	0.0E+00	00 E	X 00+30	0.0E+00 X94628.1	0.0E+00 M55582.1	0.0E+00/	0.0E+00	0.0E+00	0.0E+00 X92841.1	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 D63562.1
Expression Signal	1.4	76.0	8.0	7.78	0.95	1.66	4	1.69	1.69	1.08	3.22	0.89	0.82	1.44	1.97	1.18	2.26	1.05	8.64	1.2	1.61	1.33	24.91	2.43	2.46	5.48	5.48	0.86	12.01	0.95
ORF SEQ ID NO:	24416	24417	24419	24424	24428	24429	24430	24432	24433	24438	24437	24447	24458	24460	24463	24464	24466	24467	24469	24471	24472		24477		24480	23791	23792	24484	24490	24493
Exan SEQ ID NO:	14629	14631	14633	14637	14641	14842	14842	14844	14844	14647	14648	14860	14871	14673	14676	14877	14679	14680	14682	14684	14685	14686	14690	14692	14693	14013	14013	14697	14706	14709
Probs SEQ ID NO:	4744	4746	4748	4762	4756	4767	4757	4759	4759	4762	4763	4778	4786	4788	4791	4782	4794	4795	4797	4799	4800	4801	4806	4808	4809	4812	4812	4814	4824	4827

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Top Hit Descriptor	Homo sapiens farnesyl diphosphate synthase (farnesyl pyrophosphate synthetase, dimethylallytranstransferase, geranyltranstransferase) (FDPS) mRNA	Homo sapiens sialytransferase 8 (alpha-N-acetyneuraminate: alpha-2,8-sialytransferase, GD3 synthase) (SIAT8) mRNA	Human mRNA for transcription factor AREB6, complete cds	Human mRNA for transcription factor AREB6, complete cds	Homo sapians DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes, complete ods)	Homo sapiens chromosome 21 segment HS21C084	UI-H-BI3-alv-f-02-0-UI.s1 NCI_CGAP_Sub5 Homo sapiens cDNA clone IMAGE:3068691 3'	Homo sapiens hypothetical protein FLJ11190(FLJ1190), mRNA	Homo sapiens titin (TTN) mRNA	Homo sapiens titin (TTN) gene, alternative splice products, partial cds	Homo sapiens titin (TTN) gene, alternative splice products, partial cds	Homo sapiens titin (TTN) mRNA	Human ribosomal protein L21 mRNA, complete cds	601303729F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3638118 5'	Homo sapiens desmoplakin (DPI, DPII) (DSP) mRNA	Homo saplens mRNA for KIAA1043 protein, partial cds	Homo sapiens hypothetical protein FLJ20477 (FLJ20477), mRNA	Homo sapiens hypothetical protein FLJ20477 (FLJ20477), mRNA	Homo sapiens chromosome Xq28 melanoma antigan family A2a (MAGEA2A), melanoma antigen family A12 (MAGEA12), melanoma antigen family A2 (MAGEA2B), melanoma antigen family A3 (MAGEA3), caltractin (CALT). NAD(P)H dehydrogenase-like protein (NSDHL), and L1>	Homo sapiens chromosome Xq28 melanoma antigen family A2a (MAGEA2A), melanoma antigen family A12 (MAGEA12), melanoma antigen family A2b (MAGEA2B), melanoma antigen family A3 (MAGEA3), caltractin	(CALT), NAD(P)H dehydrogenase-like protein (NSDHL), and LI>	Homo sapiens titin (TTN) mRNA	Homo sapiens titin (TTN) mRNA	Homo sapiens titin (TTN) mRNA	Homo sapiens titin (TTN) mRNA	Homo sapiens E2F transcription factor 2 (E2F2) mRNA	Homo sapiens E6-AP ublquitin-protein ligase (UBE3A) gene, exon 3	Homo sapiens chromosome 21 segment HS2/IC009
Top Hit Database Source	IN	LΝ	FZ	NT	Ę	IN	EST_HUMAN	NT	TN	NT	TN	ĮŽ.	NT	EST_HUMAN	IN	TN	N	NT	L L		NT	NT	IN	IN	TN	NT	TN	TN
Top Hit Acession No.	4503684 NT	4506952 NT	D15050.1	D15050.1	AB026898.1	AL163284.2	AW452728.1	8922826 NT	4507720 NT	AF058332.1	AF058332.1	4507720 NT	U14967.1	BE408863.1	4758199 NT	AB028966.1	8923441 NT	8923441 NT	U82671.2		U82671.2	4507720 NT	4507720 NT	4607720 NT	4507720 NT	4758225 NT	AF016705.1	AL163209.2
Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00/		0.0E+00				0.0E+00	0.0E+00	0.0E+00	0.05+00.1	·	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00/
Expression Signal	1.52	1.06	1.31	1.31	0.86	1.34	1.45	1.2	7.99	2.81	2.81	2.95	4.34	2.58	5.37	0.39	1.66	1.66	90.		1.06	5.81	5.81	3.51	7.78	1.17	1.35	1.33
ORF SEQ ID NO:	24498	24105	24508		24515	24530	24536	24542	24490	24544	24546	24552			24569	24574	24584	24585	24596		24697	23791	23792	24602	24603		24619	
Exan SEQ ID NO:	14713	14319	14726	14726	14734	14751	14759	14766	14708	14768	14768	14774	14777	14789	14794	14804	14817	14817	14830		14830	14013	14013	14834	14835	14843	14853	14862
Probe SEQ ID NO:	4831	4837	4845	4845	4854	4871	4879	4885	4886	4888	4888	4894	4897	4910	4915	4925	4839	4939	4963		4963	4957	4957	4928	4960	4968	4978	4987

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Single Exon Probes Expressed in Heart

Top Hit Descriptor	Homo sapiens gammma-cytoplasmic actin (ACTGP3) pseudogene	Homo sapiens partial TTN gene for titin	Homo sapians titin (TTN) mRNA	Homo sapiens titin (TTN) mRNA	Homo saplens titin (TTN) mRNA	Homo saplens titin (TTN) mRNA	Bacillus amyoliquefaciens sacB gene for levensucrase (EC 2.4.1.10)	Homo sapiens vascular endothelial cadherin 2 mRNA, complete ods	Homo saplens vascular endothelial cadherin 2 mRNA, complete cds	Homo sapiens cyclophilin (USA-CYP) mRNA	Homo sapiens G-protein coupled receptor (RE2), mRNA	Homo saplens titin (TTN) mRNA	Homo sapiens titn (TTN) mRNA	Homo sapiens titn (TTN) mRNA	Homo saplens titin (TTN) mRNA	Homo sapiens PR domain containing 1, with 2NF domain (PRDM1) mRNA	Human cellular fibronectin mRNA	Human cellular fibronectin mRNA	Human hereditary haemochromatosis region, histone 2A-like protein gene, hereditary haemochromatosis (HLA-H) gene, RoRet gene, and sodium phosphate transporter (NPT3) gene, complete cds	Homo sapiens titin (TTN) mRNA	Homo sapiens titin (TTN) mRNA	Homo saplens titin (TTN) mRNA	Human offactory receptor-like gene, complete cds	Human offactory receptor-like gene, complete/cds	Homo sapiens titin (TTN) mRNA	Homo sapiens titin (TTN) mRNA	Homo saplens DNA mismatch repair protein (MLH3) gene, complete cds	Homo sapiens glypican 3 (GPC3) mRNA	Escherichia coli K-12 MG1655 section 217 of 400 of the complete genome	Homo sapiens melanoma antigen, family C, 1 (MAGEC1), mRNA	Homo sapiens melanoma antigen, family C, 1 (MAGEC1), mRNA	Homo sapiens mannosidase, alpha, class 2A, member 1 (MAN2A1), mRNA
Top Hit Database Source	LΝ	NT	Z F	Ę	۲	NT	ZI ZI	FZ	NT	F	Ę	LN FN	뉟	Į.	۲	F	ΙΝ	NT	TN TN	L'A	۲	۲۸	NT	NT	ΤN	NT	ΝT	TN	IN	NT	NT	۲
Top Hit Acession No.	D50657.1	AJ277892.1	4507720 NT	4507720 NT	4507720 NT	4507720 NT	X52988.1	AF240635.1	AF240635.1	5454153 NT	6677700 NT	4507720 NT	4507720 NT	4507720 NT	4507720 NT	4557362 NT	M10905.1	M10905.1	U91328.1	4507720 NT	4507720 NT	4507720 NT	L35475.1	L35475.1	4507720 NT	4507720 NT	AF195658.1	5360213 NT	AE000327.1	4885474 NT	4885474 NT	4758697 NT
Most Similar (Top) Hit BLAST E Value	0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	_	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00
Expression Signal	37.78	2.29	3.02	4.23	2.89	2.89	2.43	1. 28.1	1.84	1.01	1.22	11.22	11.22	14.9	14.9	1.33	1.03	1.03	1.06	3.04	6.27	6.27	1.34	1.34	9:38	9.38	0.94	1.35	0.8	1.06	96.0	1.59
ORF SEQ ID NO:		24639	24840	24642	23791	23792	24660	24681	24682	24686	24698	23791	23792	23791	23792	24708	24713	24714	24715	24726	23791	23792	24743	24744	23791	23792	24765	24768	24768	24778	24793	24794
Exan SEQ ID NO:	14865	14875	14876	14878	14013	14013	14892	14909	٠	14912	14926	14013	14013	14013	14013	14935	14940	14940	14941	14950	14013	14013		14887	14013	14013	14992	14993	14997		1	15028
Probe SEQ ID NO:	4880	0009	5001	6003	5005	5005	5018	5037	2037	5040	5054	5055	9909	5056	8036	5065	6070	5070	5071	5080	9609	2088	5098	5098	2089	5099	6124	5126	5130	5140	5159	5162

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Single Exon Probes Expressed in Heart

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Top Hit Descriptor	Homo sapiens toll-like receptor 8 (TLR8) mRNA, complete cds	Homo sapiens toll-like receptor 8 (TLR8) mRNA, complete cds	Homo sapiens placental growth hormone Isoform hGH-V3 (hGH-V) mRNA, complete cds	Homo sapiens titin (TTN) mRNA	Homo sapiens titin (TTN) mRNA	Homo sapiens titin (TTN) mRNA	Homo sapiens chromosome 21 segment HS21C085	Homo sapiens titin (TTN) mRNA	Homo saplens titin (TTN) mRNA	Homo sapiens beaded filament structural protein 1, filensin (BFSP1) mRNA	Homo sapiens aconitase (ACO2) gene, nuclear gene encoding mitochondrial protein, exon 15	Homo sapiens keratin 12 (KRT12) gene, complete cds	Homo sapiens keratin 12 (KRT12) gene, complete cds	Homo saplens protocadherin alpha 13 (PCDHA13), mRNA	RC3-GN0076-310800-013-b03 GN0076 Homo septens cDNA	Homo sapiens polycystic kidney disease-like 2 protein (PKDL2) mRNA, complete cds	Homo sapiens polycystic kidney disease-like 2 protein (PKDL2) mRNA, complete cds	H.sapiens immunoglobulin heavy chain gene, variable region	H.sapiens immunoglobulin heavy chain gene, variable region	7f10c06.x1 NCI_CGAP_CLL1 Homo saplens cDNA clone IMAGE:3294250 3'	htg9a02.x1 NCI_CGAP_Lu24 Homo saplens/cDNA clone IMAGE:3165194 3' similar to SW:Y054_HUMAN P42694 HYPOTHETICAL PROTEIN KIAA0064.;	601589422F1 NIH_MGC_7 Homo sapiens ciDNA clone IMAGE:3943804 5	801589422F1 NIH_MGC_7 Homo sapiens ciDNA done IMAGE:3943804 5'	Homo sapiens eosinophil peroxidase (EPP) gene, exon 7	one8a09.y5 NCI_CGAP_Kdd5 Homo saplens CDNA clone IMAGE:1472152 5' similar to gb:M18512 IG HFAVY CHAIN PRECIPSOR V.I. REGION (HIMAN)	Homo sapiens Sp4 transcription factor (SP4), mRNA	602118928F1 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:4276264 5	601061489F1 NIH_MGC_10 Homo sapiens cDNA done IMAGE:3447839 5'	601105891F1 NIH_MGC_15 Homo sepiens cDNA done IMAGE:2988310 5	602071372F1 NCI_CGAP_Bm64 Hamo sapiens cDNA clone IMAGE:4214272 6'	602071372F1 NCI_CGAP_Bm64 Homo sapiens cDNA clone IMAGE:4214272 5'	Homo sapians Bloom syndrome (BLM) mRNA
Top Hit Database Source	NT	Ę	N	Ę	Ŋ	NT	닏	TN	IN	FZ	IN	TN	N-I	۲	EST_HUMAN	LN	LN	IN	IN	EST_HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	LN	EST HIMAN	ŇT	EST HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	LU
Top Hit Acession No.	4F245703.1	AF245703.1	AF006061.1	4507720 NT	4507720 NT	4507720 NT	AL.163285.2	4507720 NT	4507720 NT	4502398 NT	4F093093.1	4F137286.1	AF137286.1	9256579 NT	BE931080.1	AF182034.1	AF182034.1	X56163.1	X56163.1	BE675498.1	3E220753.1	3E794412.1	3E794412.1	M29908.1	41791363 1	11421038 NT	3F665962.1	BE538857.1	BE292784.1	BF526328.1	BF526328.1	4557364 NT
Most Similar (Top) Hit BLAST E Value			0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00					0.0E+00			0.0E+00	0.0E+00	0 0F+00	'	-	_	=	0.0E+00	ı	0.0E+00
Expression Signet	1.12	1.12	1.72	10.43	10.43	6.75	1.38	3.97	3.97	98'0	15.68	2.25	2.26	2.99	3.82	3.12	3.12	1.82	1.92	5.8	1.77	1.67	1.67	5.46	181	5.42	2.91	1.92	1.31	1.8	1.8	1.71
ORF SEQ ID NO:	24797	24798	24814	23791	23792			24822	24823	24824		24830	24831	24917	24928	24930	24931	24940	24941	25032	25033	25034	25035	25037	25038	25044		25052	25075	25080	25081	26122
Exon SEQ ID NO:	15031	15031	15050	14013		15052	<u> </u>	15058	15058	15059	15086	15137		15150	15158		15162		15168	15228	15229		l	l	15234			15247	L.	<u> </u>		15987
Probe SEQ ID NO:	5165	5165	5186	5187	5187	5189	5193	5195	5185	5196	5208	5214	6214	5228	5234	5238	5238	5245	5245	5307	5308	5309	5309	5311	5313	5319	5324	5327	5333	5337	5337	5348

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Probe	Exan	ORFSEQ	Expression	Most Similar (Top) Hit	Top Hit Acession	Top Hit	Top Hit
SEQ EQ SO: -	SEQ ID	ΘΝΘ	Signal	BLAST E	o'N	Database Source	Top Hit Descriptor
5352	15272	25101	5.24	0.0E+00	AF257737.1	LN	Homo sapiens ciliary dynein heavy chain 9 (DNAH9) mRNA, complete cds
5352	15272	25102	5.24	0.0E+00	AF257737.1	LN	Homo sapiens cillary dynein heavy chain 9 (DNAH9) mRNA, complete cds
5362	15282	25114	1.45	0.0E+00	D26535.1	INT	Human gene for dihydrolipoamide succinyltransferase, complete cds (exon 1-15)
5362			1.45	0.0E+00	D26535.1	LN	Human gene for dihydrolipoamide succinyltransferase, complete cds (exon 1-15)
5374		25141	1.67	0.0E+00	11420819 NT	LN	Homo sapiens olfactory receptor, family 2, subfamily F, member 1 (OR2F1), mRNA
5386	15305		3.26	0.0E+00	BF529931.1	EST_HUMAN	602042322F1 NCI_CGAP_Brn67 Homo sapiens cDNA clone IMAGE:4179988 5'
5386			3.26	0.0E+00	BF529931.1	EST_HUMAN	602042322F1 NCI_CGAP_Brn67 Homo sapiens cDNA clone IMAGE:4179988 5
5389	15308	25161	2.35	0.0E+00	BF313139.1	EST_HUMAN	601897658F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:4126815 5'
5396	15315	25362	4.21	0.0E+00	11434392 NT	LN	Homo sapiens calclum channel, voltage-dependent, alpha 1G subunit (CACNA1G), mRNA
5420	16341		2.43	0.0E+00	AW867316.1	EST_HUMAN	MR0-SN0037-030400-001-h07 SN0037 Homo sapiens cDNA
5431	15351	25405	3.05	0.0E+00	BE292889.1	EST_HUMAN	601105291F1 NIH_MGC_15 Homo sapiens cDNA clone IMAGE:2987903 5'
5431	15351	25406	3.05	0.0E+00	BE292889.1	EST_HUMAN	601105291F1 NIH_MGC_15 Homo saplens cDNA clone IMAGE:2987903 5'
5444	ı	25420	1.31	0.0E+00	11420819 NT	TN	Homo sapiens olfactory receptor, family 2, subfamily F, member 1 (OR2F1), mRNA
24	15365	25421	1.31	0.0E+00	11420B19 NT	LN	Homo sapiens olfactory receptor, family 2, subfamily F, member 1 (OR2F1), mRNA
5451	15372	25429	5.35	0.0E+00	AF064254.1	TN	Homo sapiens very long-chain acyl-CoA synthetase homolog 1 mRNA, complete cds
5451	15372	25430	5.35	0.0E+00	AF064254.1	NT	Homo sapiens very long-chain acyl-CoA synthetase homolog 1 mRNA, complete cds
5455			2.9	0.0E+00	AJ224639.1	LN	Homo sapiens Surf-5 and Surf-6 genes
5485	15376		2.9	0.0E+00	AJ224639.1	LN	Homo sapiens Surf-5 and Surf-6 genes
5470	15390	25453	69.69	0.0E+00	M85719.1	EST_HUMAN	EST02238 Fetal brain, Stratagene (cat#936206) Homo sapiens cDNA clone HFBCM48
5474	15394	25459	3.67	0.0E+00	AW405472.1	EST_HUMAN	UI-HF-BI_0-adh-d-02-0-UI.r1 NIH_MGC_37 Homo sapiens cDNA clone IMAGE:3061658 5
5489	15408	25471	5.74		AW361877.1	EST_HUMAN	PM3-CT0263-091289-007-h05 CT0263 Homo sapiens cDNA
5489	15408	25472	5.74	0.0E+00	AW361877.1	EST_HUMAN	PM3-CT0263-091289-007-h05 CT0263 Homo sepiens cDNA
5489	15408	25473	6.74	0.0E+00	AW361877.1	EST_HUMAN	PM3-CT0263-091289-007-h05 CT0263 Homo sapiens cDNA
5491	15410	25475	1.88	0.0E+00	U36261.1	LN	Human beta-prime-adaptin (BAM22) gene, exon 13
	<u> </u>			1			zp95b11.r1 Stratagene muscle 937209 Homoleapiens cDNA clone IMAGE:627933 5' similar to gb:X03740
6531	ı	İ	3.26		AA195905.1	ESI_HUMAN	MTOSIN HEAVT CHAIN, SNELE IAL MUSICLE (HUMAN);
5532	15449		1.5		AJ006345.1	ż	Homo saplens KVLQ11 gene
5532			1.5		AJ006345.1	N	Homo saplens KVLQT1 gene!
5637		25524	2	0.0E+00	Al207616.1	EST_HUMAN	HA2981 Human fetal liver cDNA library Homo saplens cDNA
5548	15464	25534	3.98		11416801 NT	ΙN	Homo sapiens protocadherin beta 2 (PCDHB2), mRNA
5565	15471	25542	6.78	0.0E+00	BE560082.1	EST_HUMAN	601345141F1 NIH_MGC_B Homo sapiens cDNA clone IMAGE:3877843 5'
5556	15472	25543	1.58	0.0E+00	10048478 NT	. LN	Mus musculus aczonin (Acz), mRNA
5557	15473	25544	3.03	0.0E+00	U86961.1	L V	Human L-type calcium channel beta-1 subunit (CACNLB1) gene, exon 13B and isoform beta-1B, complete cds
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		_	_	-		_		_		_	_		_	_		_		<u>17 - 4.</u>	T . E .	' '	<u> </u>	11111	_	H	~/	<u> 5.</u>	11 41.	, (1 H	Ç) #:
Top Hit Descriptor	Human L-type calcium channel beta-1 subunit (CACNLB1) gene, exon 13B and Isoform beta-1B, complete cds	602036272F1 NCI_CGAP_Bm64 Homo sapiens cDNA clone IMAGE:4184321 5'	601104462F1 NIH_MGC_14 Homo sapiens cDNA clone IMAGE:3347463 5'	602185852F1 NIH_MGC_45 Homo sapiens cDNA clone IMAGE:4310076 5'	Homo saplens cadherin 20 (CDH20) mRNA, complete cds	RC5-ET0027-210600-022-G10 ET0027 Homo sapiens cDNA	601645287F1 NIH_MGC_56 Homo sepiens cDNA clone IMAGE:3930453 5'	801558060F1 NIH_MGC_58 Homo sapiens cDNA clone IMAGE:3827775 5'	601558060F1 NIH_MGC_58 Homo sapiens cDNA clone IMAGE:3827775 5'	2008h06.r1 Soares_parathyroid_tumor_NbHPA Homo sapiens cDNA clone IMAGE:321755 5'	2008h06.r1 Soares_parathyroid_tumor_NbHPA Homo sapiens cDNA clone IMAGE:321755 5'	Homo saplens familial mental retardation protein 2 (FMR2) gene, exon 14	601158515F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3505323 5'	601512630F1 NIH_MGC_71 Homo saplens cDNA clone IMAGE:3914238 5'	Hamo sapiens KIAA0735 gene product, synaptic vesicle protein 2B hamolog (KIAA0735), mRNA	Homo sapiens KIAA0735 gene product; synaptic vesicle protein 2B homolog (KIAA0735), mRNA	Homo saplens potassium voltage-gated channel, Shal-related subfamily, member 2 (KCND2), mRNA	740h01.11 Scares_NhHMPu_S1 Home saplens cDNA clone IMAGE:665905 5' similar to SW:YY05_HUMAN P42694 HYPOTHETICAL MYELOID CELL LINE PROTEIN 5.	240h01.r1 Scares_NhHMPu_S1 Homo sapieris cDNA clone IMAGE:865905 5' similar to SW:YY05_HUMAN P42694 HYPOTHETICAL MYELOID CELL LINE PROTEIN 5.	Human T cell surface glycoprotein CD-6 mRNA, complete ods	Human T cell surface glycoprotein CD-6 mRNA, complete cds	AU137772 PLACE1 Homo sapiens cDNA clone PLACE1007201 5'	Human G protein-coupled receptor GPR-9-8 gene, complete cds	zq81d03.r1 Stratagene hNT neuron (#937233) Homo sapiens cDNA clone IMAGE:648005 5' similar to TR:G854195 G854195 LEUKOCYTE SURFACE PROTEIN :	Homo sapiens xylosyltransferase II (XT2), mRNA	Homo sapiens xylosyltransferase II (XT2), mRNA	601109532F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3350622 5'	Human anion exchanger (AE1) gene, exons 1-20	Homo sapiens peptide transporter 3 (LOC51286), mRNA
Top Hit Detabase Source	F	EST_HUMAN	EST_HUMAN	EST_HUMAN	TN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	EST HUMAN	EST_HUMAN	۲	IN	LN	EST HUMAN	EST_HUMAN	N FN	FN	EST_HUMAN	LN	EST HUMAN	LN	LN	EST HUMAN	NT	LN
Top Hit Acession No.	J86961.1	3F338835.1	BE273983.1	3F569905.1	0.0E+00 AF217289.1	3E828144.1	3E958636.1	3F031742.1	3F031742.1	W33069.1	N33069.1	4F012618.1	3E280197.1	3E889610.1	11433071 NT	11433071 NT	9789886 NT	0.0E+00 AA193508.1	0.0E+00 AA193506.1	J34625.1	J34625.1	U137772.1	45982.1	0.0E+00 AA204740.1	11645913 NT	11545913 NT	E2571		11435830 NT
Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00 B	0.0E+00 B	0.0E+00	0.0E+00 B	0.0E+00	0.0E+00 B	0.0E+00	0.0E+00	0.0E+00 W	0.0E+00	0.0E+00 B	0.0E+00 B	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 A	0.0E+00 U	0.0E+00	0.0E+00	0.0E+00	0.0E+00 B	0.0E+00	0.0E+00
Expression Signal	3.03	2.1	2.83	1.74	2.47	1.89	1.41	1.66	1.66	1.54	1.54	2.18	3.57	2.74	. 1.63	1.63	10.66	1.29	1.29	13.03	13.03	1.41	3.4	4.14	3.57	3.57	2.8	1.47	1.38
ORF SEQ ID NO:	25545	25553	25555					25613	25614		25634		_	25642	25654	25655	25668	25671	25672	26690	L		26758	25770	25771	25772	25790		25811
Exan SEQ ID NO:	15473	15480		15489	15510	16511	15515	15530	15530	ı		15546	15548	15551	15561	15561	19450	15574	15674	15589	15589		15649	15663	15684	ı	ı		15701
Probe SEQ ID NO:	2557	5564	9955	5574	9839	5597	5601	5615	5615	5631	5631	5632	5634	5638	5649	5649	0999	5663	5863	5680	2880	5730	5741	99.29	5756	5756	5775	5784	5795

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	_	_	_	Τ-	_	_	г-	_	_	_	_	_	T	_		_		-82	· 1	Ψ.	-	-				'n	بوك	4	إسالا	بالبا	11 IT	71 11
Top Hit Descriptor	AV650020 GLC Homo sapiens cDNA clone GLCCAD09 3'	UI-HF-BL0-acc-g-12-0-UI.s1 NIH_MGC_37 Homo sapiens cDNA clone IMAGE:3058751 3'	y27b03.r1 Scares placenta Nb2HP Homo sapiens cDNA clone IMAGE:149933 5'	Human gene for the light and heavy chains of myeloperoxidase	601305368F1 NIH_MGC_39 Homo sapiens cDNA clone IMAGE:3639616 5'	601305368F1 NIH_MGC_39 Homo sapiens cDNA clone IMAGE:3639616 5'	AU119245 HEMBA1 Homo sapiens CDNA clone HEMBA1005360 5'	AU119245 HEMBA1 Homo sepiens cDNA clone HEMBA1005360 5	ws25c07.x1 NCI_CGAP_GC6 Homo saplens,cDNA clone IMAGE:2498220 3'	601105344F1 NIH_MGC_15 Homo sapiens cDNA clone IMAGE:2987963 5'	601105344F1 NIH_MGC_15 Homo sapiens cDNA clone IMAGE:2987963 6'	UI-HF-BL0-aco-h-02-0-UI.r1 NIH_MGC_37 Homo sapiens cDNA done IMAGE:3059931 5	UI-HF-BLO-acc-h-02-0-UI.r1 NIH_MGC_37 Homo sapiens cDNA clone IMAGE:3059931 5'	AV719444 GLC Homo sapiens cDNA clone GLCEHC06 5'	Homo sapiens low voltage-activated T-type calcium channel alpha 1G splice variant CavT.1a (CACNA1G)	mRNA, complete cds	au96h08.y1 Schneider fetal brain 00004 Horno sapiens cDNA clone IMAGE:2784159 6' similar to TR:015390 015390 GT24, [3] TR:043840 TR:043206 ;	aus6th08.y/ Schneider fetal brain 00004 Homo sapiens cDNA clone IMAGE:2784159 5' similar to TR:015380 015390 GT24. [3] TR:043840 TR:043208:	601587561F1 NIH MGC 7 Homo seplens cDNA clone IMAGE:3941847 5'	601512058F1 NIH_MGC_71 Homo saplens cDNA clone IMAGE:3913311 5	601512058F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3913311 5'	Human antigen CD27 gene, exons 1-2	Homo saplens chromosome 21 segment HS21C004	Hamo sapiens chromosome 21 segment HS2; C004	Homo sapiens zona peliucida glycoprotein 3A (sperm receptor) (ZP3A), mRNA	#31f11.x1 NCI_CGAP_GC8 Home saplens cDNA clone IMAGE:2242413 3' similar to SW:WNT3_MOUSE	FIGURE AND INSTITUTE THE PROJECT TO SELVENCE AND SELVENCE	ronto sapiens Zinc iniger nomeodomem Protein (A I DF 1-A) mixtyA, comprete cas	zw52c03.r1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA clone IMAGE:773668 5'	QV3-BN0047-300800-278-c06 BN0047 Homo sapiens cDNA	AU125928 NT2RM4 Homo sapiens cDNA clone NT2RM4002430 6"	PM3-HT0520-230200-002-c08 HT0520 Honc sapiens cDNA
Top Hit Database Source	EST HUMAN	EST_HUMAN	EST_HUMAN	Ä	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN		NT	EST_HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	N _T	NT	NT	₽N	1000	ES - HUMAN	Z		EST_HUMAN	EST_HUMAN	EST_HUMAN
Top Hit Acession No.	AV650020.1	0.0E+00 AW575598.1	0.0E+00 H01255.1	0.0E+00 X15377.1	0.0E+00 BE735989.1	0.0E+00 BE735989.1	0.0E+00 AU119245.1	AU119245.1	AI989483.1	0.0E+00 BE293153.1	0.0E+00 BE293153.1	0.0E+00 AW 406348.1	0.0E+00 AW 406348.1	0.0E+00 AV719444.1		AF190860.1	0.0E+00 AW163640.1	0 0E+00 AW163640 1	0.0E+00 BE799873.1	0.0E+00 BE889813.1	0.0E+00 BE889813.1	24493.1	0.0E+00 AL163204.2	0.0E+00 AL163204.2	TN 6865009			32832.1	0.0E+00 AA434584.1	0.0E+00 BE925875.1	0.0E+00 AU126928.1	BE169131.1
Most Similar (Top) Hit BLAST E Value	0.0E+00 A	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 A	0.0E+00	0.0E+00	0.0E+00					· 0.0E+00 A	0.0E+00	001-100	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	200	0.0E+00 A	0.0E+00 L	0.0E+00	0.0E+00	0.0E+00	0.0E+00
Expression Signal	8.1	2.81	4.21	1.99	4.25	4.25	11.67	11.57	1.56	4.81	4.81	1.36	1.36	1.69		2.24	3.45	3.45	4 79	7.23	7.23	3.71	2.15	2.15	3.54		4.13	A).	3.58	1.48	1.44	7.44
ORF SEQ ID NO:		25848	25850	25857	25862	25863	25869	25870		25882	25883	25931	25932	25951		25958	25964	25965			26982	25988	25991	26992	25999				l			26076
SEQ ID NO:	15732	15736	15738	Į į	•	l	15763	15753		15765	15765	15807	15807	15826		15835	15841	15841	1	١.	l	15866	15869	15869	15875		_ [_			15896		15944
Probe SEQ ID NO:	5826	5830	5832	5838	5843	5843	5847	5847	5854	5859	5859	5891	5994	1269 1692 1		5930	9999	9265	5951	5955	5965	5961	5964	5864	5970		2 82	97/4	5980	5991	6011	6041

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601443867F1 NIH_MGC_65 Homo sapiens ¢DNA clone IMAGE:3847697 5'
7b49f03.x1 NCI_CGAP_Lu24 Homo sapiens|cDNA clone IMAGE:3231581 3' similar to SW:GC95_HUMAN Q08379 GOLGIN-95; 1549f03 x1 NCI_CGAP_Lu24 Homo sapiensicDNA clone IMAGE:3231581 3' similar to SW:GG95_HUMAN Homo sapiens catenin (cadherin-associated protein), delta 2 (neural plakophilin-related arm-repeat protein) Human chromosome 16 creatine transporter (SLC6A8) and (CDM) paralogous genes, complete cds EST366876 MAGE resequences, MAGC Horio sapiens cDNA 601113958F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3354566 5 zp88e03.r1 Stratagene HeLa cell s3 937216 Homo sapiens cDNA clone IMAGE:627292 5' H.sapiens mRNA for latent transforming growth factor-beta binding protein (LTBP-2) H.sapians mRNA for latent transforming growth factor-bata binding protein (LTBP-2) #34g03.r1 Soares_NhHMPu_S1 Homo sapiens cDNA clone IMAGE:665332 601339977F1 NIH_MGC_53 Homo sapiens cDNA clone IMAGE:3682267 5' 601443867F1 NIH_MGC_65 Homo sapiens cDNA clone IMAGE:3847697 5' 602185852F1 NIH_MGC_45 Homo sapiens cDNA clone IMAGE:4310076 5' 602185852F1 NIH_MGC_45 Homo sapiens cDNA clone IMAGE:4310076 5' 601148954F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:3501829 5' Human type IV sodium channel alpha polypepiide (SCN4A) gene, exon 19 601889823F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:4123948 Homo saplens ciliary dynein heavy chain 9 (DNAH9) mRNA, complete cds Homo sapiens ciliary dynein heavy chain 9 (DNAH9) mRNA, complete cds Homo sapiens melanoma antigen, family B, 2 (MAGEB2), mRNA Top Hit Descriptor CM1-HT0877-060900-397-g11 HT0877 Homo sapiens cDNA L5-GN0032-180900-145-d07 GN0032 Homo sepiens cDNA L3-ST0024-230799-001-B01 ST0024 Homo sapiens cDNA L3-ST0024-230799-001-B01 ST0024 Homo sapiens cDNA Homo sapiens cadherin 20 (CDH20) mRNA, complete cds Homo sapiens cadherin 20 (CDH20) mRNA, complete cds Homo sapiens NALP1 mRNA, complete cds Homo sapiens CD6 antigen (CD6), mRNA Human MYCL 2 gene, complete cds Q08379 GOLGIN-95. (CTNND2), mRNA EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN NT EST_HUMAN HUMAN HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN 11034810 NT 05.1 EST HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN Top Hit Database Source 눋 늘 되되보 눋 z 눋 11435626 **Top Hit Acession** 0.0E+00 U41302.1 0.0E+00 AW954806.1 0.0E+00 BE254103.1 0.0E+00 BE550162.1 0.0E+00 BF088376.1 0.0E+00 AA195106.1 0.0E+00 BE566381.1 0.0E+00 BE566381.1 0.0E+00 BE867889.1 0.0E+00 AA180755.1 0.0E+00 AI940621.1 0.0E+00 AI940621.1 0.0E+00 1142 0.0E+00 BE262941.1 0.0E+00 237976.1 0.0E+00 Z37976.1 0.0E+00 AF257737.1 BF569905.1 BE262941.1 0.0E+00 BE550162.1 0.0E+00 AF217289.1 BF569905.1 0.0E+00 AF257737. 0.0E+00 AF310105. BF306996. 0.0E+00 BE867889. 0.0E+00 J03069.1 0.0E+00 0.0E+00 0.0E+00 0.0E+00 **Most Similar** (Top) Hit BLAST E Value 3.07 2.75 5.63 2.32 3.15 6.08 6.08 13.15 10.28 1.78 13.15 2.07 2.07 1.46 3.84 2.57 2.32 2.83 2.18 3.68 5.61 1.34 1.41 £ Expression Signal 26201 26202 26208 24867 24876 24842 24844 24846 24846 24846 24851 28150 28153 26118 26213 26227 26242 26191 24872 28130 26131 26158 ORF SEQ 26157 24873 ÖNO SEQ ID NO: 15945 16048 16054 16078 16054 16059 15109 15995 5995 16012 15973 15982 15990 16019 15112 15125 15128 15126 16064 16067 16075 15104 15109 16019 15127 15127 16091 16092 6094 6094 6118 6154 6168 6169 6170 6183 8228 Probe SEQ ID 6063 6071 6071 609 6039 6121 6126 6135 6142 6146 6146 6169 6170 8175 6178 6180 6225 6101 9101 6181

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Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
6237	16103	26253	6.9	0.0E+00	AU133213.1	EST_HUMAN	AU133213 NT2RP4 Homo sapiens cDNA clane NT2RP4001556 6
6253	16119		2.44	0.0E+00	AU143708.1	EST_HUMAN	AU143706 Y79AA1 Homo sapiens cDNA clone Y79AA1002365 5'
6260	16126		1.31	0.0E+00	BE891286.1	EST_HUMAN	601431819F1 NIH_MGC_72 Homo sapiens cDNA done IMAGE:3917164 5'
6260			1.31	0.0E+00	1	EST_HUMAN	601431819F1 NIH_MGC_72 Homo sapiens cDNA done IMAGE:3917164 5'
6273			1.97	0.0E+00	AF137286.1	NT	Homo saplens keratin 12 (KRT12) gene, complete cds
6273	15137	24831	1.97	0.0E+00	AF137286.1	NT	Homo sapians keratin 12 (KRT12) gene, complete cds
6286	16150		3.63	0.0E+00	11436899 NT	LN.	Homo sapiens vitamin D (1,25- dihydroxyvitamin D3) receptor (VDR), mRNA
6286	16150	26306	3.63	0.0E+00	11436699 NT	NT	Homo sapiens vitamin D (1,25- dihydroxyvitaniin D3) receptor (VDR), mRNA
							qc67a07.x1 Scares, placenta_8to9weeks_2NbHP8to9W Homo sapians cDNA_clone IMAGE:1714644 3' similar to SW:ARSD_HUMAN P61689 ARYLSULFATASE D PRECURSOR ;contains element HGR
6302	16166	26323	25.51	0.0E+00	Al128344.1	EST_HUMAN	repetitive element;
							qc67a07.x1 Soares_placenta_8to9weeks_2NbHP8to9W Homo sapiens cDNA clone IMAGE:1714644.3' similar to SW:ARSD_HUMAN P51689 ARYLSULFATASE D PRECURSOR contains element HGR
8302	16166	26324	25.51	0.0E+00	AI128344.1	EST_HUMAN	repetitive element;
6304	ı	26326	18.73	0.0E+00	11426392 NT	NT	Homo sapiens myosin, heavy polypeptide 8, skeletal muscle, perinatal (MYH8), mRNA
6304				0.0E+00	11426392 NT	NT	Homo saplens myosin, heavy polypeptide 8, skeletal muscle, perinatal (MYH8), mRNA
6306	L		14.06	0.0E+00	BF337375.1	EST_HUMAN	602035089F1 NCI_CGAP_Brn64 Homo sapions cDNA clone IMAGE:4182839 5'
							zn60f09.r1 Stratagene muscle 937209 Homo saplens cDNA clone IMAGE:562601 5' similar to TR:G806562
8308	- 1		5.1	0.0E+00	AA128453.1	EST_HUMAN	G806562 NEBULIN.;
6328		26353	6.72	0.0E+00	AF005213.1	NT	Homo sapiens ankyrin 1 (ANK1) mRNA, complete cds
6328			8.72	0.0E+00	AF005213.1	IN	Homo sapiens ankyrin 1 (ANK1) mRNA, complete cds
6337		26360	7.55	0.0E+00	X70172.1	IN	H. sapiens DNA for ZNGP2 pseudogene, exon 4
6338		26362	11:09	0.0E+00	U45448.1	TN	Human P2x1 receptor mRNA, complete cds
6339	16202	26363	11.09	0.0E+00	U45448.1	NT	Human P2x1 receptor mRNA, complete cds
6347	16210		1.43	0.0E+00	AW956503.1	EST_HUMAN	EST368573 MAGE resequences, MAGD Homo saplens cDNA
6349	16212	26374	2.54	0.0E+00	AW950516.1	EST_HUMAN	EST362586 MAGE resequences, MAGA Horrio sapiens cDNA
8380	18281	26411	1 87	001000	1 9CEOEC MV	NVW IT LOD	xb39a05.yf NCI_CGAP_Lu31 Homo sapiens cDNA clone IMAGE 2578840 5' similar to TR:Q08050 Q08050 HNRSH TRANSCRIPTION FACTOR GENESIS
8400			. A	00E+00	A11417553 4	EST HIMAN	AU117553 HFMB41 Homo saniens cDNA clone HFMB41001881 5'
6404		26422	3.64	0.0E+00	11427135	.1	Homo sapians glucagon-like peptide 2 receptor (GLP2R), mRNA
6411	<u> </u>		54.65	0.0E+00	AA2116	EST HUMAN	zn56702.r1 Stratagene muscle 937209 Homo sapiens cDNA clone IMAGE:562203 5' similar to gb:X03740 MYOSIN HEAVY CHAIN, SKELETAL MUSCLE (HUMAN);
970	ł		·				Topic of Microsoft Direction of Date of State of
2407	16321	20480	4.20	0.0E+00	A1/52561.1	EST HOMAIN	CITTOODAXI NOTIFIE DELIGITATION DE CEUS DOTO SEPTETS COUNTY CIONE IN LIDO-CITTOODA

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Table 4
Single Exon Probes Expressed in Heart

Homo sapiens sema domain, seven thrombospondin repeats (type 1 and type 1-like), transmembrane domain zi81b04.r1 Stratagene schizo brain S11 Homo sepiens cDNA done IMAGE:728719 5' similar to TR:G30048. G300482 POL=REVERSE TRANSCRIPTASE HOMOLOG (RETROVIRAL ELEMENT); domo sapiens dynactin 1 (DCTN1) gene, alteinatively spliced products, exons 7 through 32 and complete Homo sapiens dynactin 1 (DCTN1) gene, alternatively spliced products, exons 7 through 32 and complete cn17d05.xt Normal Human Trabecular Bone Cells Homo sapiens cDNA clone NHTBC_cn17d05 random au93b08.x1 Schneider fetal brain 00004 Homo sapiens cDNA clone IMAGE:2783799 3' similar to DKFZp761P092_r1 761 (synonym: hamy2) Homo sapiens cDNA clone DKFZp761P092 5 DKFZp761P092_r1 761 (synonym: hamy2) Homo sapiens cDNA clone DKFZp761P092 5 601485254F1 NIH_MGC_69 Homo sapiens cDNA clone IMAGE;3887773 5 JI-HF-BN0-akj-f-01-0-UI.r1 NIH_MGC_50 Homo saplens cDNA clone IMAGE:3077496 5' Homo saplens ATP-binding cassette, sub-family A (ABC1), member 3 (ABCA3), mRNA 601481713F1 NIH_MGC_68 Homo sapiens ciDNA clone IMAGE:3884258 5' 601481713F1 NIH_MGC_68 Homo sapiens ciDNA clone IMAGE:3884258 5' 2001c06.rl Stratagene colon (#937204) Homo|sapiens ciDNA clone IMAGE:566410 5' TR:060463 060463 TYPE-2 PHOSPHATIDIC ACID PHOSPHOHYDROLASE. [1] 601885465F1 NIH_MGC_57 Homo sapiens cONA clone IMAGE:4103729 5 601305658F1 NIH_MGC_39 Homo sapiens cDNA clone IMAGE:3639903 5' 602163008F1 NIH_MGC_81 Homo saplens cDNA clone IMAGE:4294128 5' 601593156F1 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:3947365 5'
AU120424 HEMBB1 Homo sapiens cDNA clone HEMBB1000665 5' 601578195F1 NIH_MGC_9 Hamo saplens cDNA clone IMAGE:3926998 5' 601578195F1 NIH_MGC_9 Hamo saplens cDNA clone IMAGE:3926998 5' (TM) and short cytoplesmic domain, (semaphorin) 54 (SEMA5A), mRNA Homo saplens transient receptor potential chainnel 5 (TRPC5), mRNA 601593156F1 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:3947 AU120424 HEMBB1 Homo sapiens cDNA clone HEMBB1000655 5' AU129622 NT2RP2 Homo sapiens cDNA clone NT2RP2005913 5 AU142402 Y79AA1 Homo saplens cDNA clone Y79AA1000277 6" Fop Hit Descriptor Human amyloid-beta protein (APP) gene, exon 11 Human amyloki-beta protein (APP) gene, exon 111 쁑 EST HUMAN EST HUMAN EST HUMAN EST_HUMAN EST HUMAN **EST HUMAN** EST_HUMAN HUMAN EST_HUMAN EST HUMAN HUMAN Top Hit Database Source HUMAN EST_HUMAN EST_HUMAN EST HUMAN EST HUMAN EST_HUMAN EST HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST EST Ę 눋 11417342|NT 4501848 NT 6912735 **Fop Hit Acesslon** 0.0E+00 BE877693.1 0.0E+00 AW500549.1 0.0E+00 AW157233.1 0.0E+00 AF064205.1 0.0E+00 BE739870.1 0.0E+00 AU120424.1 0.0E+00 AI752561.1 0.0E+00|AF064205.1 0.0E+00 BF217905.1 BE739870.1 0.0E+00 AA397551.1 0.0E+00 AU142402.1 0.0E+00 BE745597.1 0.0E+00 BE745597.1 BE787610.1 0.0E+00 BE736046.1 AL120124.1 0.0E+00 AU129622. 0.0E+00 AL120124.1 ġ 0.0E+00 BE787610. AA149791. 0.0E+00 BF673098.1 AU120424. 0.0E+00 M34872.1 0.0E+00 M34872.1 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 (Top) Hit BLAST E dost Simila Value 6.49 4.25 8 83 2.98 4.97 60.88 60.88 1.18 5.37 1.52 1.52 8.73 1.98 14.35 1.16 3.72 3.97 7.54 1.96 1.35 1.31 Expression Signal 26487 26530 28578 28588 26590 26592 26593 26614 26813 ORF SEQ 26531 26571 26615 26729 26749 26750 26783 26810 26842 26843 26691 26784 ΘŃΟ SEQ ID 16358 16358 16525 16321 16374 16389 16392 16413 16413 16414 16414 16502 16594 16594 16626 16432 16534 16654 16610 16821 16654 16397 16557 16568 16654 16534 ġ Probe SEQ ID 6462 6489 6439 6515 6534 6539 6550 6555 6555 6556 6556 6574 6574 6645 6674 6677 6714 6742 6747 6530 6714 8776 8778 6654 6854 6688 6730 ġ

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Onigo Excit Tobas Expressed in Teal.	Top Hit Descriptor	Homo sapiens Xq pseudoautosomal region; segment 1/2	7d76s04.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3278862.3' similar to TR:095793 095793 STAUFEN PROTEIN.;	wi60b10.x1 NCI_CGAP_Bm25 Home sapiens cDNA clone INAGE:2429275 3' similar to SW:COGT_HUMAN P55281 MATRIX METALL OPROTEINASE-14 PRECURSOR	601334760F1 NIH MGC 39 Homo septens cDNA clone IMAGE:3688655 5	601334780F1 NIH_MGC_39 Homo saplens cDNA clone IMAGE:3688655 5	Homo saplens Chediak-Higashi syndrome 1 (CHS1), mRNA	Homo sapiens Chediak-Higashi syndrome 1 (CHS1), mRNA	273a08.s1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:727958.3' similar to gb:S85655 PROHIBITIN (HUMAN);	QV3-D10045-221299-046-c07 DT0045 Homo saptens cDNA	QV3-DT0045-221299-048-c07 DT0045 Horno saplens cDNA	601452412F1 NIH MGC 88 Homo saplens cDNA clone IMAGE:3856179 5	601452412F1 NIH_MGC_68 Homo capiens cDNA clone IMAGE:3856179 5'	Homo sapiens chromosome 21 segment HS2/C009	Homo saplens chromosome 21 segment HS21 C009	601431238F1 NIH_MGC_72 Homo sapiens cDNA clone IMAGE:3916569 5'	Homo sapiens mitogen-activated protein kinase kinase kinase 13 (MAP3K13), mRNA	Homo sapiens mitogen-activated protein kinase kinase kinase 13 (MAP3K13), mRNA	H.sapiens mRNA for gamma-glutamytransferase	H.saplens mRNA for gamma-glutamytransferase	H.sapiens mRNA for gamma-glutamytransferase	xo46601.x1 NCI_CGAP_Ut1 Homo sapiens cDNA clone IMAGE:2707032.3' similar to gb:M14123_cds4 RETROVIRUS-RELATED POL POLYPROTEIN (HUMAN);	HUM084C02B Clontech human fetal brain poly/A+ mRNA (#6535) Homo sapiens cDNA clone GEN-084C02		601236488F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3608709 5	z32e04.r1 Soares ovary fumor NbHOT Homo sapiens cDNA clone IMAGE:724062 5:	601900571F1 NIH_MGC_19 Homo sapiens cONA clone IMAGE:4129744 5'	UI-H-BI1-adr-e-12-0-UI.s1 NCI_CGAP_Sub3 Home sapiens oDNA clone IMAGE:2717687 3'	4GE:2717687 3'		
פופי	Top Hit Database Source	LN	EST_HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	INT	TN	EST_HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	N	TN	EST_HUMAN	INT	INT	NT	IN	NT	EST_HUMAN		EST HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN
5	Top Hit Acession No.	AJ271735.1	BE674157.1	AI885671.1		BE563650.1	11427235 NT	11427235 NT	AA398511.1	AW364874.1	AW364874.1	BE612586.1	BE612586.1	AL163209.2	AL163209.2	BE890797.1	4768695 NT	4758695 NT	X98922.1	X98922.1	X98922.1	AW513513.1		D52650.1	BE378495.1	AA410545.1	BF313946.1	AW 139673.1	AW139673.1	BE260272.1	BF700165.1
	Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0€+00	0.0E+00	0.0E+00	0.0E+00
	Expression Signal	2.72	2.2	1.36	1.31	1.31	1.44	1.44	3.89	1.45	1.45	1.21	1.21	1.25	1.25	2.01	2.4	2.4	2.85	2.85	2.85	1.36		3.64	4.46	1.31	4.32	1.41	1.41	2.39	1.83
	ORF SEQ ID NO:	26852	26878	26879	26887	26888	26892	26898		82692	26929	26942	26943	26856	26957		26984	26985	27026	27027	27028			27083	27081	27083		27088	27089	27104	27106
	Exen SEQ ID NO:	16662	16689	16690	16696		16703	16703	16730	16735	16735	16748	16748	16758	16758	16778	16791		- 1		16833	16870		109/2	16888	16892	16893	16898	16898	16915	16917
	Probe SEQ ID NO:	6783	6810	6811	6817	6817	6824	6824	6851	6856	6856	6869	6889	6879	6879	6889	6913	6913	6955	6955	6955	6993	-	2	7011	7915	7016	7021	7021	2038	7040

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Table 4
Single Exon Probes Expressed in Heart

	Т	Т	Т	Т	Т	Т	Т	_	Т	Т	Т	Т	Т	Т	T	Т	٢	<u>"</u>	Τ"	T	<u>, , , , , , , , , , , , , , , , , , , </u>	"	T"	Γ	<u> </u>	Z	4	413	115	N 15.	31 31;
Top Hit Descriptor	602127664F1 NIH_MGC_58 Homo sapiens cDNA clone IMAGE:4284542 5	602127664F1 NIH_MGC_56 Homo sapiens aDNA clone IMAGE:4284542 5'	ক80g02.s1 NCI_CGAP_Lu5 Homo sapiens ≎DNA clone IMAGE:1802194 3' similar to gb:M36072.60S RIBOSOMAL PROTEIN L7A (HIMAN):	Homo sapiens ankyrin 1, erythrocytic (ANK1), transcript variant 1, mRNA	Homo sapiens ankyrin 1, erythrocytic (ANK1), transcript variant 1, mRNA	Homo sapiens ITGB4 gene for Integrin beta 4 subunit, exons 3-41	AV718377 FHTB Hamo sepiens cDNA clone FHTBAAF11 5	xw73c07xf NCI_OGAP_Pan1 Homo sapiens cDNA clone IMAGE:2833644 3' similar to gb:X53587 INTEGRIN BETA-4 SUBUNIT PRECURSOR (HUMAN):	AU124051 NT2RM2 Homo sapiens cDNA clone NT2RM2001575 5'	Homo sapiens mRNA for KIAA0454 protein, partial cds	hf48a09.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2935096 3'	hf48a09.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2935096 3'	DKFZp434C1814_s1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434C1814 3'	DKFZp434C1814_s1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434C1814 3'	Homo saplens killer inhibitory receptor 2-2-1 (KIRZZ1) and killer inhibitory receptor 2-2-2 (KIR2Z2) genes, partial cds	Homo saplens mRNA for KIAA1512 protein, partial cds	Homo sapiens fumor protein p73 (TP73), mRIVA	Human Ig rearranged H-chain epsilon-3 pseudogene, constant region	Homo saplens mRNA for KIAA0823 protein, partial cds	Homo saplens mRNA for KIAA0823 protein, partial cds	AV660739 GLC Homo sepiens cDNA clone GLCGKG123'	Homo sapiens polycystin-L (PKDL), mRNA	601141119F1 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:3140740 5'	601141119F1 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:3740740 5'	Human mRNA for GABA-A receptor, alpha 1 subunit	wq34a12.x1 NCI_CGAP_GC6 Homo sapiens cDNA clone IMAGE:2473150 3' similar to SW:MGB3_HUMAN	O15480 MELANOMA-ASSOCIATED ANTIGEN B3;	Homo sapiens protocedherin alpha 8 (PCDHA3), mRNA	EST370381 MAGE resequences, MAGE Homo sapiens cDNA	Human endogenous retrovirus, complete genoine	Homo saplens MAP-kinase activating death domain (MADD), mRNA
Top Hit Database Source	EST_HUMAN	EST_HUMAN	EST HIMAN	NT	NT	NT	EST_HUMAN	EST HUMAN	EST HUMAN	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	TN	IN	TN	TN	LN	NT	EST_HUMAN	NT	EST HUMAN	EST_HUMAN	ΤN		EST_HUMAN	N	EST_HUMAN	NT	NT
Top Hit Acession No.	BF700165.1	BF700165.1	AA982527 1	10947037 NT	10947037 NT	Y11107.3	AV718377.1	AW337277.1			AW592233.1	AW592233.1	AL040428.1	AL040428.1	AF133901.1	AB040945.1	11422857 NT	K01241.1		0.0E+00 AB020830.1	0.0E+00 AV660739.1	7706638 NT	E315402.1	5315402.1	14766.1		195460	9256595	W958311.1	9635487 NT	11436995 NT
Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.05+00	0.05+00	0.05+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 A	0.0E+00	0.0E+00	0.0E+00	0.0E+00			0.0E+00	0.0E+00	0.0E+00	0.0E+00 B	0.0E+00 BI	0.0E+00		0.0E+00 A	0.0E+00	0.0E+00 A	0.0E+00	0.0E+00
Expression Signal	1.83	1.83	6.35	3.54	3.54	1.28	1.45	3.64	1.57	2.64	4.41	4.41	2.84	2.84	1.17	18.6	3.97	1.25	2.65	2.65	1.96	3.43	3.86	3.86	1.91		2.12	4.49	1.54	1.49	6.88
ORF SEQ ID NO:	27107		27137					27169		27218				27242	27243			27268			27277				27313		_				27352
Exan SEQ ID NO:	16917	16917	16946	16950	16950	16965	16972	16976	16979	17024	17025	17025	17053	17053	17054	17055	17075	17081	17084	17084	17087	17090	17108	17108	17118		- [17131	ĺ	17148	17157
Probe SEQ ID NO:	7040	7040	7069	7073	7073	7088	7095	7099	7102	7147	7148	7148	7176	7176	7117	7178	7198	7204	7207	7207	7210	7213	7231	7231	7241		R _V	7254	7263	7269	7280

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	Top Hit Descriptor	Homo sapiens mRNA for KIAA0578 protein, partial cds	601589294F1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:3943463 5'	601510247F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3911986 5	601510247F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3911986 5'	EST50505 Gall bladder I Homo saplens cDNA 5' end	EST50505 Gall bladder i Homo sapiens cDNA 5' end	ba09f05.y1 NIH_MGC_7 Home saplens cDNA clone IMAGE:2823873 5' similar to gb:L35049 Mus musculus Bcl-xL mRNA, complete cds (MOUSE):	ba09f05.y1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:2823873 5' similar to gb:L35049 Mus musculus	Bcl-xL mRNA, complete cds (MOUSE);	602023150F1 NCI_CGAP_Brn67 Homo sepiens cDNA clone IMAGE:4158300 5	QV2-HT0698-250700-282-b08 HT0698 Homo sapiens cDNA	DKFZp434L0120_r1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434L0120 5	ow60h01.x1 Soares_NSF_F8_9W_OT_PA_F_S1 Homo sepiens cDNA clone IMAGE:1651249.3' similar to TR-O14677 O14677 KIAA0171 PROTEIN	Home control brockedied College and Educate Profess (El 19960A) with	Domo sapiens hypothetical CZDZ zinc linger protein PLJZZSO4 (PLJZZSO4), mixivA Domo sapiens himshatical (ODO sinc fines ringle) El 103604 (El 103604), mBNA	The same of the sa	qm09a08.x1 NG_CGAP_Lu5 Homo sapiens cDNA clone IMAGE:1881298 3' similar to SW:RL2B_HUMAN P28316 60S RIBOSOMAL PROTEIN L23A. ;	qm09a05.x1 NCI_CGAP_Lu5 Homo saplens cDNA clone IMAGE:1881298 3' similar to SW:RL2B_HUMAN_P23316 60S RIBOSOMAL PROTEIN L23A.	EST386026 MAGE resequences, MAGC Homo saplens cDNA	Homo sapiens polycystic kidney disease 2-like protein (PKD2L) gene, exon 8	601109942F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3350722 5'	601466828F1 NIH_MGC_67 Homo sapiens dDNA clone IMAGE:3870007 5'	601486828F1 NIH_MGC_67 Homo sapiens clONA clone IMAGE:3870007 5	eu86c04.y1 Schnelder fetal brain 00004 Homo sapiens cDNA clone IMAGE:2783142 5' similar to gb:M36072 60S RIBOSOMAL PROTEIN L7A (HUMAN);	601145054F2 NIH_MGC_19 Hamo sapiens cDNA clone IMAGE:3160477 5'	C06158 Human pancreatic Islat Homo sapiens cDNA clone hbc5605	C06158 Human pancreatic islet Homo sapiens cDNA clone hbc5605	601578683F1 NIH_MGC_9 Homo saplens cDNA clane IMAGE:3927548 5'	Homo saplens solute carrier family 21 (organic anion transporter), member 9 (SLC21A9), mRNA	Homo sapiens solute carrier family 21 (organic anion transporter), member 9 (SLC21A9), mRNA
	Top Hit Database Source	N	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	l	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HIMAN	LICE COLUMN	Z	-	EST_HUMAN	HST HIMAN	EST HUMAN	N	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	LN	LN
	Top Hit Acession No.	AB011160.1	BE794823.1	BE883843.1	BE883843.1	AA344601.1	AA344601.1	BE207063.1		BE207063.1	BF348013.1	BE712515.1	AL042278.1	AIDBBDA3 1	745504F	TIN 1200151	10100011	A1290909.1	A1290909 1	AW953836.1	AF153466.1	BE255829.1	BE781382.1	BE781382.1	AW 163779.1	BE263191.1	C06158.1	C08158.1	BE746215.1	11437282 NT	11437282 NT
	Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00			0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00	00+30 O	00+300	20.10	0.05+00	2	0.0E+00	0.05+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00
	Expression Signal	1.44	2.58	1.24	1.24	1.8	1.6	1.38		1.38	2.71	8	11.81	1 27	190 0	2.00	3	8.86	8 8	9.1	3.92	6.4	1.37	1.37	7.21	2.85	3.98	3.98	3.22	1.93	1.93
	ORF SEQ ID NO:	27365	27368	27374	27376	27383		27428		27427	27551	27457		אבעש		97007		26632	55996	L	27464		27529	27530	27531	27541		27563			27575
	Exan SEQ ID NO:	17168	17167	17174	17174	17184	17184	17227		17227	17346	17252	17273	17202	2077	10442	<u> </u>	16444	16444	16443	17259	17321	17323	17323	17324	17335	17358	17358	17360	17369	17369
	Probe SEQ ID NO:	7290	7291	7298	7298	7308	2308	2360		7360	7368	7383	7406	7425	247	2007	1	7431	7431	7432	7450	7461	7463	7463	7464	7475	7488	7488	7490	7499	7499

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Onigio Lyon Flories Expressed III hear	Top Hit Descriptor	Homo saciens solute carrier family 21 (organic anion transporter), member 9 (SLC21A9), mRNA	801873425F1 NIH MGC 21 Homo sapiens DNA clone IMAGE:3956238 5	Homo saplens keratin 2e (KRT2E) gene, complete cds	Homo sapiens keratin 2e (KRT2E) gene, complete cds	RC2-BT0642-130300-017-g01 BT0642 Homb sapiens oDNA	UI-HF-BN0-ekg-b-12-0-UI.r1 NIH MGC 50 Homo sapiens cDNA clone IMAGE:3076943 5'	UI-HF-BN0-akg-b-12-0-UI.r1 NIH_MGC_50 Homo sapiens cDNA clone IMAGE:3076943 5'	Homo sapiens chromosome 9 duplication of the T cell receptor beta locus and tryosinoxen gene families	Homo sapiens chromosome 9 duplication of the T cell receptor beta locus and truncinonen nene families	ULHF-BNO-aki-c-07-0-ULT NIH MGC 50 Homo seciens cDNA clone IMAGE:3077384 5	Multiple sclerosis associated retrovirus polyprotein (pol) mRNA, partial cds	AIGF=androgen-induced growth factor AIGF [human, placenta, Genomic/mRNA, 498 nt. segment 5 of 5]	AIGF=androosen-induced growth factor AIGF (numan placenta Genomic/mRNA 408 nt seament 5 of হা	601334603F1 NIH MGC 39 Homo sapiens cDNA clone IMAGE:3688680 5	CM2-CT0311-301189-043-h11 CT0311 Homo saplens cDNA	AU132349 NT2RP3 Homo saplens cDNA clone NT2RP3004260 5	AU132349 NT2RP3 Homo sapiens cDNA clone NT2RP3004260 6'	601595558F1 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:3949383 5'	601595558F1 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:3949383 5'	Homo sapiens KIAA0345 gene product (KIAA0345), mRNA	AU132349 NT2RP3 Hamo sapiens cDNA clone NT2RP3004260 5'	Homo sapiens protocadharín alpha 12 (PCDH alpha12) mRNA, complete cds	Homo sapiens leucocyte immunoglobulin-like receptor-1 mRNA, complete cds	Homo sapiens leucocyte immunoglobulin-like receptor-1 mRNA, complete cds	MR4-TN0114-110900-101-e04 TN0114 Homo saplens cDNA	601155227F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3138798 5'	xn72b01.x1 NCI_CGAP_CML1 Homo saplens cDNA clone IMAGE:2699977 3' sImiler to gb:X02152_cds1 L-LACTATE DEHYDROGENASE M CHAIN /HIMANN:	Homo saplens Chediak Higashi syndrome 1 (CHS1), mRNA	AU143873 Y79AA1 Homo sapiens cDNA clone Y79AA1002307 5
אַטון זין ווטאַן פונ	Top Hit Database Source	LN	EST_HUMAN	E	F	EST HUMAN	EST HUMAN	EST_HUMAN	LΝ	Ę	EST HUMAN	LZ LZ	TN	IN	EST HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	ΤZ	EST_HUMAN	NT	LN.	NT	EST_HUMAN	EST_HUMAN	EST HUMAN	トラ	T HUMAN
	Top Hit Acession No.	11437282 NT	BE900549.1	AF019084.1	I٩	BE082977.1	0.0E+00 AW 500293.1	0.0E+00 AW500293.1	F029308.1	F029308.1		0.0E+00 AF009668.1		0.0E+00 S78466.1	0.0E+00 BE563320.1		Γ			0.0E+00 BE740490.1	7662067 NT	U132349.1		0.0E+00 AF009220.1			0.0E+00 BE280793.1		7235	0.0E+00 AU143673.1
	Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 A	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.05+00
	Expression Signal	1.93	1.47	2.59	2.59	1.47	1.76	1.76	1,25	1.25	2.45	1.34	2.56	2.56	2.57	1.62	2.17	2.17	7.73	. 7.73	1.78	2.22	1.86	. 2.72	2.72	1.65	2.44	1.74	1.91	5.98
	ORF SEQ ID NO:	27576		27591	27692		27626	27627	27629	27630	27645		27691	27692		27701	27722			1	ı	27756	27757	27776	27777	27784	27785	27800	27810	27826
	Exon SEQ ID NO:		17302	17381	17381		17410	17410	17414		17431	17458	17472	17472			17500	17500	17509	17509	17516	17532	17533	17551	17551	17558	17570	17578	17586	17603
	Probe SEQ ID NO:	7499	7514	7530	7530	7548	7559	7559	7583	7563	7580	7607	7621	7621	7622	7630	7650	7650	7659	7659	7666	7682	7683	701	7701	7708	7728	7728	7736	7753

Single Exon Probes Expressed in Heart Page 405 of 413 Table 4

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a74b01.x1 NCI_CGAP_Kid8 Homo sapiens cDNA clone IMAGE:2807401 3' similar to gb:M69086 MOESIN zq06h11.r1 Stratagene muscle 937209 Homo saplens cDNA clone IMAGE:628965 5' similar to TR:G407097 nab45e12.x1 Soares_NSF_F8_9W_OT_PA_F_S1 Homo saplens cDNA clone IMAGE:32652713 231f01.r1 Soares_pregnant_uterus_NbHPU Homo sapiens cDNA clone IMAGE:503545 57 231f01.r1 Soares_pregnant_uterus_NbHPU Homo sapiens cDNA clone IMAGE:503545 57 2997h11.r1 Stratagene muscle 937209 Homo saplens cDNA clone IMAGE:628197 5 Homo sapiens partial RANBP7 gene for RanEP7/Importin7 and partial ZNF143 gene Homo sapiens partial RANBP7 gene for RanBP7/Importin7 and partial ZNF143 gene 602037045F1 NCI_CGAP_Bm84 Homo sapiens cDNA clone IMAGE:4184939 55 602037045F1 NCI_CGAP_Bm84 Homo sapiens cDNA clone IMAGE:4184939 55 601570712F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3845403 5 1078764F1 NIH_MGC_12 Homo sapiens cDNA clone IMAGE:3464703 5' 601467419F1 NIH_MGC_67 Homo sapiens cDNA clone IMAGE:3870700 5 601570712F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3845403 5' Homo sapiens hypothetical C2H2 zinc finger protein FLJ22504 (FLJ22504), 601573895F1 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:3835198 5 601573895F1 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:3835198 5 FB23A4 Fetal brain, Stratagene Homo sapiens cDNA clone FB23A4 3'end Homo sapiens killer cell inhibitory receptor KIRCI gene, exons 2, 3, and 4 Homo sapiens mRNA for actin binding protein/ABP620, complete cds AU136637 PLACE1 Homo saplens cDNA clone PLACE1004737 5 AU136637 PLACE1 Homo sapiens cDNA clone PLACE1004737 5 AU143673 Y79AA1 Homo sepiens cDNA clone Y79AA1002307 Top Hit Descriptor Homo sapiens mRNA for estrogen receptor beta, complete cds Homo sapiens mRNA for estrogen receptor beta, complete cds RC2-BT0642-150200-012-d03 BT0642 Homo sapiens cDNA RC2-BT0642-150200-012-d03 BT0642 Homo sapiens cDNA AV654765 GLC Homo sepiens cDNA clone GLCDZC07 3' AV711075 Cu Homo sapiens cDNA clone CuAAKG05 5' Homo sapiens KIF4 (KIF4) mRNA, complete cds Homo sapiens HEF like Protein (HEFL), mRNA Homo sapiens HEF like Protein (HEFL), mRNA G407097 165KD PROTEIN. (HUMAN); EST_HUMAN EST_HUMAN EST HUMAN EST HUMAN EST HUMAN EST_HUMAN EST_HUMAN EST_HUMAN HUMAN HUMAN EST HUMAN EST HUMAN EST_HUMAN HUMAN **EST HUMAN** EST HUMAN **EST HUMAN** EST HUMAN EST_HUMAN EST_HUMAN Top Hit Database Source EST_HUMAN EST HUMAN EST EST 눋 Ę ż 11421001 **Fop Hit Acession** 0.0E+00 AA194770.1 0.0E+00 AJ295844.1 0.0E+00 AJ295844.1 0.0E+00 AA196387.1 0.0E+00 AW517960.1 0.0E+00 BE743215.1 0.0E+00 BE743215.1 0.0E+00 AV711075.1 0.0E+00 AU136637.1 0.0E+00 BE730772.1 0.0E+00 BE730772.1 0.0E+00]AB006590.1 AU136637.1 0.0E+00 AA131248.1 0.0E+00 AA131248.1 AB029290.1 0.0E+00 BE549213.1 0.0E+00 BE781742.1 BE082720.1 0.0E+00 AF17930B.1 AF072408.1 0.0E+00 AB006590.1 0.0E+00 BF340331.1 0.0E+00 BF340331.1 BF436218.1 0.0E+00 AV654765.1 0.0E+00 BE082720.1 0.0E+00 BE082720.1 ģ 0.0E+00 T03078.1 0.0E+00 BF436218 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 Aost Similar (Top) Hit BLAST E Value 5.43 3.45 5.19 2.23 1.69 5.98 7.52 2.48 2.48 2.96 2.13 2.13 4.01 1.17 1.46 1.24 1.64 3.27 5.43 1.37 3.55 6.06 1.65 1.69 2.33 2.05 Expression Signal 27833 27915 27916 27868 27869 27892 27895 28036 28065 27879 27937 27960 27885 27987 28072 28092 28112 ORF SEQ 27993 27994 28111 27991 ÖΝΩ SEQ ID 17608 17608 17845 17645 17654 17673 17754 17865 17924 17635 17873 17796 17832 17851 17858 17865 17692 17742 1753 17753 17823 17834 ö Probe SEQ ID 7758 7804 7865 7865 7865 8015 7823 7823 7892 7905 7905 7946 7972 7982 7984 8009 8008 8015 7795 7904

7903 7993 7973

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Table 4
Single Exon Probes Expressed in Heart

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Top Hit Descriptor	AV711075 Cu Homo sapiens cDNA clone CuAAKG05 5'	RC3-ST0197-120200-015-a03 ST0197 Homo saplens cDNA	EST375636 MAGE resequences, MAGH Homo sapiens cDNA	Homo sepiens ATP-binding cassette, sub-family A (ABC1), member 3 (ABCA3), mRNA	Homo sapiens ATP-binding cassette, sub-family A (ABC1), member 3 (ABCA3), mRNA	wy61f09.xt Soares_NSF_F8_9W_OT_PA_P_S1 Homo sapiens cDNA clone IMAGE:2553065.3' similar to TR:060566 Q80568 VDX:	TCAAP3D0917 Pediatric acute myelogenous leukemia cell (FAB M1) Baylor-HGSC project=TCAA Homo sapiens cDNA clone TCAAP0917	wb28a12.x1 NCI_CGAP_GC8 Homo saplens cDNA clone IMAGE:2306974 3' similar to contains element MSR1 MSR1 repetitive element:	wb28a12.x1 NCI_CGAP_GC6 Homo capiens/cDNA clone IMAGE:2306974.3' similar to contains element MSR1 MSR1 repetitive element	Homo sapiens NOD2 protein (NOD2), mRNA	Homo saplens NOD2 protein (NOD2), mRNA	UI-HF-BL0-acm-d-04-0-UI.r1 NIH_MGC_37 Hamo sapiens cDNA clone IMAGE:3059383 5'	Homo sapiens hypothetical protein FLJ20078 (FLJ20079), mRNA	Homo sapiens 5-hydroxytryptamine (serotonin) receptor 1E (HTR1E) mRNA	Homo sapiens 6-hydroxytryptamine (serotonin) receptor 1E (HTR1E) mRNA	wu32b08.x1 Soares_Dieckgraefe_colon_NHCD Homo sapiens cDNA clone IMAGE:2521715 31	601505204F2 NIH_MGC_71 Hamo sapiens cDNA clone IMAGE:3906865 5'	601434522F1 NIH_MGC_72 Hamo sapiens cIDNA clane IMAGE:3919636 5'	Homo saplens myosin, heavy polypeptide 2, skeletal muscle, adult (MYH2), mRNA	Homo sapiens myosin, heavy polypeptide 2, skeletal muscle, edult (MYH2), mRNA	601874332F1 NIH_MGC_21 Hamo saplens cDNA clone IMAGE:3957343 5'	zp95b11.r1 Stratagene muscle 937209 Homo saplens cDNA clone IMAGE:627933 5' similar to gb:X03740 MYOSIN HEAVY CHAIN. SKELETAL MUSCIE (HUMAN).	601588829F1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:3943015 6'	AV727362 HTC Homo seplens cDNA clone HTCAQH08 5'	AV727362 HTC Homo sepiens cDNA clone HTCAQH06 5'	xy04g10 x1 NCI_CGAP_Lym12 Homo saplens cDNA clone IMAGE:2852228 3' similar to gb:M60854 40S RIBOSOMAL PROTEIN S16 (HUMAN):	AU135741 PLACE1 Homo sepiens CDNA dans PLACE1002794 5'	
Top Hit Database Source	EST_HUMAN	EST HUMAN	EST HUMAN	NT	LN	EST_HUMAN	EST_HUMAN	EST HUMAN	EST HIMAN	1.	F	EST_HUMAN	1	F	N	EST HUMAN	EST_HUMAN	EST_HUMAN	۲N	۲N	EST_HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	
Top Hit Acession No.	4V711075.1	0.0E+00 AW813783.1	4W963563.1	0.0E+00 11431124 NT	11431124 NT	4W057621.1	0.0E+00 BE243270.1	A1852239.1	52239.1	11545911	11545911 NT	1404795.1	11424829 NT	4504536 NT	04536		0.0E+00 BE882109.1	3E891630.1	8923939 NT	3939	0.0E+00 BE903304.1	195905.1		Γ			0.0E+00 AU135741.1	
Most Similar (Top) Hit BLAST E Value	0.0E+00 AV	0.05+00	0.0E+00,		0.0E+00	0.0E+00 AW	0.0E+00 E	0.0E+00 AIB	0 0E+00 AI6	0.0E+00	0.0E+00	0.0E+00 AW	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 E	0.0E+00 E	0.0E+00	0.0E+00	0.0E+00 E	0.0E+00 AA	0.0E+00 E	0.0E+00	0.0E+00	0.0E+00 AW	0.0E+00 A	
Expression Signal	2.33	6.11	6.43	2.5	2.5	1.89	1.92	4.86	4.86	2.91	2.91	2.01	4.8	9.16	9.16	2.73	3.04	10.56	22.38	22.38	1.91	4.05	4.69	6.8	6.8	17.98	217	
ORF SEQ ID NO:	28171		28178	28191	28192	28195	28200	28201	28202	28209	28210			28225	28226	28227	28231	28233	28234	28235	28247	25516	28269	28277	28278	28296	28301	
Exen SEQ ID NO:	17924		17931	17942	17942	17946	17950	17951	17951	1	17959	, ,	17975	17976	17976	17971	17980	17984	17986	17986	18000	15448	18022	18031	18031	18044	18049	
Probe SEQ ID NO:	8032	8034	8040	8051	8051	8054	8039	8060	8060	8088	8088	8081	8084	888	8085	9808	88	888	88	88	<u>8</u>	8113	8134	8143	8143	8156	8161	

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kn72b01.x1 NCI_CGAP_CML1 Homo sapiens cDNA clone IMAGE:2699977 3' similar to gb:X02152_cds1 Lkw66f01.x1 NCI_CGAP_Pan1 Homo sapiens cDNA clone IMAGE.2832985 3' similar to gb:X17115 IG MU hg13d02.x1 Soares_NFL_T_GBC_S1 Home saplens cDNA clone IMAGE:2945475 3' similar to contains ng13d02.x1 Soares_NFL_T_GBC_S1 Homd sapiens cDNA clone IMAGE:29454753' similar to contains hg13d02x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2945475 3' cimilar to contains CHAIN C REGION (HUMAN); UI-H-BI3-alh-a-01-0-UI.s1 NCI_CGAP_Sub5 Homo sepiens cDNA clone IMAGE:2736649 3' UI-H-BI3-alh-a-01-0-UI.s1 NCI_CGAP_Sub5 Homo sepiens cDNA clone IMAGE:2736649 3' 602037014F1 NCI_CGAP_Brn64 Homo sapiens cDNA ctone IMAGE:4184979 5 qf43c03.x1 Soares_tests_NHT Homo sapien; cDNA clone IMAGE:1752772.37 qf43c03.x1 Soares_tests_NHT Homo sapien; cDNA clone IMAGE:1752772.37 HSC3IC031 normalized Infant brain cDNA Homo sapiens cDNA clone c-3lc03 601119248F1 NIH_MGC_17 Homo saplens cDNA clone IMAGE:3029219 5' 601148357F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:3163310 5' Human protein kinase C substrate 80K-H (PRKCSH) gene, exon 15-17 602141405F1 NIH_MGC_46 Homo sapiens cDNA clone IMAGE: 4302432 5 601582046F1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:3936539 5' AU116908 HEMBA1 Homo saplens cDNA clone HEMBA1000255 5 AU118386 HEMBA1 Homo saplens cDNA clone HEMBA1003486 5 Top Hit Descriptor QV0-UM0093-170400-191-d06 UM0093 Homp sepiens cDNA QV0-UM0093-170400-191-d06 UM0093 Homb sapiens cDNA QV4-ST0234-121199-032-b06 ST0234 Homo sapiens cDNA Homo saplens of cardlac alpha-myosin heavy chain gene Homo saplens of cardiac alpha-myosin heavy chain gene Homo sapiens mRNA for KIAA0545 protein, partial cds Homo saplens mRNA for KIAA0587 protein, partial cds LACTATE DEHYDROGENASE M CHAIN (HUMAN) Homo saplens ribosomal protein L31 (RPL31) mRN/ Homo sapiens RGH1 gene, retrovirus-like element Homo sapiens insulin receptor (INSR), mRNA H.sapiens mRNA for H1 histamine receptor element MSR1 repetitive element; element MSR1 repetitive element element MSR1 repetitive element EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN NT EST_HUMAN EST_HUMAN EST HUMAN EST_HUMAN EST_HUMAN HUMAN EST HUMAN EST HUMAN EST HUMAN HUMAN EST_HUMAN Top Hit Database EST_HUMAN **EST HUMAN** EST_HUMAN HUMAN Source EST. 눋 눋 11424726 Top Hit Acession 0.0E+00 AW 59333.1 0.0E+00 AW 59333.1 0.0E+00 AW 59333.1 0.0E+00 AW338094.1 0.0E+00 AW451230.1 0.0E+00 AW 451230.1 0.0E+00 AW804516.1 0.0E+00 AW804516.1 0.0E+00 AW 236269.1 0.0E+00 AB014567.1 0.0E+00 BE298449.1 0.0E+00 AB011117.1 0.0E+00 BE792155.1 0.0E+00 BF684061.1 0.0E+00 AU118386.1 0.0E+00 AW391837.1 0.0E+00 BF340308.1 ġ 0.0E+00 BE261209.1 0.0E+00 U50328.1 AI149809.1 0.0E+00 Z34897.1 0.0E+00 F13069.1 0.0E+00 D10083.1 0.0E+00 Z20656.1 0.0E+00 | Z20858.1 0.0E+00 0.0E+00 0.0E+00 (Top) Hit BLAST E 0.0E+00 **fost Similar** Value 2.03 3.45 3.45 2.12 2.92 5.64 2.35 1.88 59.52 3.47 3.05 4.62 18.83 5.64 6.16 6.77 6.77 49.5 2.53 68.7 Expression Signal 28304 28305 28308 28308 28313 28328 28346 28359 28363 28379 28379 28394 28394 28395 28403 28330 28380 28386 28392 ORF SEQ 28371 28374 ÖΝΩ 18054 18056 18078 SEQ ID 18054 18054 18057 18078 10179 18125 18130 18142 18145 18162 18064 18077 18092 18109 18120 18121 18124 18151 18152 18153 18081 18106 18131 18151 18161 Š ÿ 8166 8166 SEQ ID 8166 8168 8169 8176 8192 8194 8196 8208 8224 8227 8240 8241 8244 8245 8251 8265 8272 8273 8282 8283 8262 819 8250 8271 8271 ö

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Table 4
Single Exon Probes Expressed in Heart

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Τορ Hit Descriptor	be04407.01 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:2823373 5' similar to TR:076022 076022 E1B- 55KDA-ASSOCIATED PROTEIN .	ba04d07.y1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:2823373 5' similar to TR:076022 O76022 E1B- 55KDA-ASSOCIATED PROTEIN ;	tal muscle (MYH4), mRNA	apiens cDNA	piens cDNA	piens cDNA	zp95b11.r1 Stratagene muscle 937209 Homo saptens cDNA clone IMAGE:627833 5' similar to gb:X03740 MYOSIN HEAVY CHAIN, SKELETAL MUSCLE (HUIMAN);	mo sapiens cDNA clone IMAGE:3085026 3'	mo saplens cDNA clone IMAGE:3085026 3'	7LACE1001381 5	A clone IMAGE:3889207 5'	A clone IMAGE:3889207 5'	A clone IMAGE:4089710 5'	al cds	al ods	i), mRNA	i), mRNA	or 5A (EIF5A) mRNA	A clone IMAGE:4289502 5'	one IMAGE:2847177 5'	sp:	wf20e11.x1 Soares_Dieckgraefe_colon_NHUC Homo sapiens cDNA done IMAGE:2351180 3: similar to dobin87789 IG GAMMA-1 CHAIN C REGION (HUMAN);	A clane IMAGE:4123948 5'	A clone IMAGE:4123948 5'	piens cDNA	A clane IMAGE:3924577 5'		A clone IMAGE:4081715 5'	ba04dp7.y1 NIH_MGC_7 Homo saplens cDNA clone IMAGE:2823373 5' similar to TR:076022 076022 E1B3 55KDA-ASSOCIATED PROTEIN.;
<u>Б</u>	ba04do7.yf NIH_MGC_7 Homo sapiens cDNA c	ba04d07.y1 NIH_MGC_7 Homo sapiens cDINA of 55KDA-ASSOCIATED PROTEIN:	Homo sapiens myosin, heavy polypeptide 4, skeletal muscle (MYH4), mRNA	QV0-UM0091-120900-385-b12 UM0091 Homo sapiens cDNA	RC3-HT0230-040500-110-h04 HT0230 Horio sapiens cDNA	RC3-HT0230-040500-110-h04 HT0230 Homo sapiens cDNA	2995b11.r1 Stratagene muscle 937209 Homb sapiens cDN MYOSIN HEAVY CHAIN, SKELETAL MUSCLE (HUMAN);	UI-H-BI4-aok-b-10-0-UI.s1 NCI_CCAP_Sub3 Homo sapiens cDNA cione IMAGE:3085026 3	UHH-BI4-ack-b-10-0-UI.s1 NCI_CGAP_Sub8 Homo saplens cDNA clone IMAGE:3085026 3	AU135170 PLACE1 Homo sapiens cDNA clone PLACE1001381 5	601486828F1 NIH_MGC_69 Homo sapiens :DNA clone IMAGE:3889207 5	601486828F1 NIH_MGC_69 Homo sapiens cDNA clone IMAGE:3889207 5'	601875830F1 NIH_MGC_55 Homo sapiens cDNA clone IMAGE:4099710 5	Homo sapiens mRNA for KIAA1316 protein, partial cds	Homo saplens mRNA for KIAA1316 protein, partial cds	Homo saplens retinoblastoma-like 2 (p130) (RBL2), mRNA	Homo sapiens retinoblastoma-like 2 (p130) (RBL2), mRNA	Homo sapiens eukaryotic translation initiation factor 5A (EIF5A) mRNA	602134132F1 NIH_MGC_81 Homo sapiens ¿DNA clone IMAGE:4289502 5	dr04g05.x1 NIH_MGC_3 Homo sapjens cDNA clone IMAGE:2847177 5'	Human gamma actin-like pseudogene, complète cds	wf20e11.x1 Soares_Dieckgraefe_colon_NHUC Homo sat qb:M87789 IG GAMMA-1 CHAIN C REGION (HUMAN):	601889823F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:4123948 5	601889823F1 NIH_MGC_17 Homo saplens cDNA clone IMAGE.4123948 5	QV2-NN0054-230800-333-e04 NN0054 Homo sapiens cDNA	601439605F1 NIH_MGC_72 Homo sapiens cDNA clone IMAGE:3924577 5	Homo saplens golgin-like protein (GLP), mRNA	601861947F1 NIH_MGC_53 Homo sapiens cDNA clone IMAGE:4081715 5	ba04d07.y1 NIH_MGC_7 Homo saplens cDNA o 55KDA-ASSOCIATED PROTEIN.;
Top Hit Database Source	EST_HUMAN	EST_HUMAN	뉟	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	M	M	M	NT	TN	EST_HUMAN	EST_HUMAN	M	EST HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	EST HUMAN	F	EST_HUMAN	EST_HUMAN
Top Hit Acession No.	BE206848.1	BE206846.1	11024711 NT	BF093687.1	BE148076.1	BE148076.1	AA195905.1	BF507876.1	BF507876.1	AU135170.1	BE876401.1	BE876401.1	BF240536.1	AB037737.1	AB037737.1	11430868 NT	11430868 NT	4503544 NT	BF576267.1	AW328173.1	M55083.1	Al660968.1	BF306996.1	BF306996.1	BF362462.1	BE897051.1	B923698 NT	BF207662.1	BE206846.1
Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00			0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00
Expression Signal	2.59	2.69	2.85	2.01	2.9	2.9	6.47	4.47	4.47	2.16	5.62	5.62	10.32	3.05	3.05	3.49	3.49	6.1	2.49	5.44	120.65	3.18	3.64	3.64	26.88	4.07	2.89	2.24	4.61
ORF SEQ ID NO:	28628	28629	28648	28651			28665	28687	28688			78697		28713		28717	28718	28734	28741	28744		28760			28758		28793		28833
Exan SEQ ID NO:	18363	18363	18383	18386	18390	18330	18398	18418	18418	18423	18427	18427	18435				18449	18463	18470	18472	18475	18479	1	l	18486	18504	18512	18514	18550
Probe SEQ ID NO:	8490	8480	8511	8514	8518	8518	8628	8546	8546	8553	8557	8557	8566	8577	8577	8581	8581	8596	8603	8605	8608	8612	8614	8614	8620	8639	8648	8650	8661

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be04d07.y1 NIH_MGC_7 Homo saplens cDNA clone IMAGE:2823373 5' similar to TR:O76022 O76022 E1B. 68KDA-ASSOCIATED PROTEIN ; 7/27/12.x1 NCI_CGAP_CLL1 Homo septens CDNA clone IMAGE:3285919 3' similar to TR:000409 000409 rh22b10.x1 NCI_CGAP_Co16 Homo saplens cDNA clone IMAGE:3316699 3' similar to TR:Q13458 Q13458 wn83g03x1 NCI_CGAP_Ut1 Homo sapiens cDNA clone IMAGE:2452468 3' similar to gb:S37431 LAMININ nl42c08.s1 NCI_CGAP_Pr4 Homo sapiens cDNA clone IMAGE:1043342 similar to gb:M95178 ALPHA-ACTININ 1, CYTOSKELETAL ISOFORM (HUMAN); vz91h01.x1 NCI_CGAP_Brn25 Hamo sapiens cDNA clone IMAGE.2566225 3' similar to WP:F53H10.2 nz11c07.s1 NCI_CGAP_GCB1 Homo sapienis cDNA clone IMAGE:1287468 3' similar to TR:Q13686 Q13686 ALKB HOMOLOG PROTEIN.; 601501090F1 NIH_MGC_70 Homo sapiens cDNA clone IMAGE:3902926 5' nz11c07.s1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:1287468 3' similar to TR:Q13686 dr02b08.x1 NIH_MGC_3 Homo sapiens cDNA clone IMAGE:2846919 5' IL5-HT0731-020500-077-f05 HT0731 Homo sapiens cDNA DKFZp434G178_71 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434G178 5' DKFZp434G178_r1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434G178 5 Homo sapiens polycystic kidney disease-associated protein (PKD1) gene, complete cds Homo sapiens polycystic kidney disease-associated protein (PKD1) gene, complete cds AU138211 PLACE1 Homo sapiens cDNA clone PLACE1008077 6' 601441096F1 NIH_MGC_72 Homo sapiens cDNA clone IMAGE:3916270 5' wp06g08x1 NCL_CGAP_Kid12 Homo sapleris cDNA clone IMAGE:2464094 3' 601572186T1 NIH_MGC_55 Homo saplens cDNA clone IMAGE:3839012 3' 601572186T1 NIH_MGC_55 Homo saplens cDNA clone IMAGE:3839012 3' AU141882 THYRO1 Homo sapiens oDNA clone THYRO1001398 5 AU141882 THYRO1 Homo saplens cDNA clone THYRO1001398 5 Top Hit Descriptor MR4-ST0118-261099-012-b03 ST0118 Homo sepiens cDNA MR4-ST0118-261099-012-b03 ST0118 Homo sepiens cDNA QV0-CT0225-101289-071-f06 CT0225 Homo saplens cDNA Homo saplens KiAA0247 gene product (KIAA0247), mRNA Q13686 ALKB HOMOLOG PROTEIN. CE11040 ZINC FINGER, C2H2 TYPE CHECKPOINT SUPPRESSOR 1. RECEPTOR (HUMAN); TRIO.: EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN **EST HUMAN** EST_HUMAN EST_HUMAN EST HUMAN EST HUMAN EST HUMAN Top Hit Database EST_HUMAN EST_HUMAN EST_HUMAN EST HUMAN EST_HUMAN EST HUMAN EST_HUMAN EST_HUMAN Source EST 11435244 NT Top Hit Acesslon 0.0E+00 AI834954.1 0.0E+00 AW327895.1 0.0E+00 BE185656.1 0.0E+00 AL046540.1 BE206846.1 0.0E+00 AW753028.1 0.0E+00 AA760913.1 0.0E+00 AW006022.1 0.0E+00 AW387776.1 0.0E+00 AW387776.1 0.0E+00|AA558707.1 AA760913.1 0.0E+00 BE676347.1 0.0E+00 L39891.1 0.0E+00|BF00233.1 BE910546.1 ģ 0.0E+00 AL046540.1 0,0E+00 AI923116.1 BE748899. 0.0E+00/AU141882. 0.0E+00 BE622317. AU141882. 0.0E+00 L39891.1 0.0E+00 AU138211. 0.0E+00 0.0E+00 0.0E+00 0.0E+00 Aost Similar (Top) Hit BLAST E Value 2.38 4.73 2.78 2.78 1.91 3.19 12.53 4.18 4.18 10.47 4.61 4.74 2.35 2.97 2.57 10.47 Expression Signal ORF SEQ ID NO: 24915 28842 28800 28812 28924 28836 28823 28863 28864 28888 28943 28962 28963 28876 28982 28869 28941 18518 SEQ ID 18552 15148 18529 18539 18580 18580 18640 18673 18557 18529 18589 18599 18611 18640 18850 18653 19474 18673 18690 18584 18650 <u>9</u> Probe SEQ ID 8663 8998 8670 8712 8784 8797 8861 8861 8878 8700 8722 8724 8724 8737 8772 8827 8840 8843 8661 888 8772 8712 8728 8827 8837 8837 ÿ

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Table 4
Single Exon Probes Expressed in Heart

	T	T	Τ	T	1	T	Т	Т	Т	T	Т	Т	T	≥	≥	T	11-	Ť	T	ŕ	ľ	Ï	T .	T	Ť		ŕ	1 −3	T	
Top Hit Descriptor	Hamo sapiens KIAA0247 gene product (KIAA0247), mRNA	Human beta-prime-adaptin (BAM22) gene, exon 5	601237691F1 NIH MGC 44 Homo saplens cONA clone IMAGE:3609623 5'	601237691F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3609623 5'	205602.r1 Stratagene muscle 937209 Homo tapiens cDNA clone IMAGE:662203 5' similar to gb:X03740 MYOSIN HEAVY CHAIN: SKELETAL MUSCLE (HUMAN):	UI-HF-BNO-ama-c-01-0-UI.r1 NIH MGC 50 Homo sapiens cDNA clone IMAGE:3081217 5'	601590568F1 NIH_MGC_7 Hamo sapiens cDNA clone IMAGE:3944708 5	601491821F1 NIH_MGC_69 Homo sapiens cDNA clone IMAGE:3894220 5'	Human von Willebrand factor pseudogene corresponding to exons 23 through 34	601299403F1 NIH_MGC_21 Homo saplens cDNA clone IMAGE:3629544 5'	Homo sapiens protein kinase, AMP-activated, tipha 2 catalytic subunit (PRKAA2) mRNA	Homo sepiens protein kinase, AMP-activated, alpha 2 catalytic subunit (PRKAA2), mRNA	Homo sapiens protein kinase, AMP-activated, aipha 2 catalytic subunit (PRKAA2), mRNA	Homo sapiens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively spliced	Homo saplens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively spiliced	602/55722F1 NIH MGC 83 Home septens CDNA clare IMAGE 4296725 51	602155722F1 NIH MGC 83 Homo saplens cDNA clane IMAGE 4296725 5'	601676357F1 NIH MGC 21 Hamo sapiens cDNA clone IMAGE:3958935 5'	601897524F1 NIH_MGC_19 Hamo sapiens cDNA clone IMAGE:4127069 5'	601897524F1 NIH_MGC_19 Homo saplens cDNA clone IMAGE:4127069 5'	Human lambda-Immunoglobulin constant region complex (germline)	Human lambda-immunoglobulin constant region complex (germline)	601890534F1 NIH_MGC_17 Homo saplens cDNA clone IMAGE:4131416 5'	RC4-NN0025-120600-016-b07 NN0025 Homo sapiens cDNA	RC4-NN0025-120600-016-b07 NN0025 Homo sapiens cDNA	601177407F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:3532968 5'	Homo saplens myosin, heavy polypeptide 1, skeletal muscle, adult (MYH1), mRNA	Homo sapiens myosin, heavy polypeptide 1, skeletal muscle, edult (MYH1), mRNA	Homo sapiens myosin, heavy polypeptide 4, skejetal muscle (MYH4), mRNA	HSB77E122 STRATAGENE Human skeletal misscle cDNA library. cat. #938215. Homo sepiens cDNA clone
Top Hit Database Source	TN	IN	EST HUMAN	HUMAN	EST HUMAN	Т	Т	T HUMAN	Г	EST_HUMAN	LN			LN FX	Ż	T HUMAN	Т	HUMAN	Г	r_HUMAN	LN			EST_HUMAN	EST_HUMAN	EST_HUMAN				
Top Hit Acesslon No.	11435244 NT	U36253.1	BE379254.1	BE379254.1	0.0E+00 AA211683.1	0.0E+00 AW 505430.1	0.0E+00 BE794758.1	0.0E+00 BE879633.1	0.0E+00 M60676.1	0.0E+00 BE409993.1	11427345	11427345 NT	11427345 NT	0.0E+00 AF223391.1	0.0E+00 AF223391.1	0.0E+00 BF681841.1	0.0E+00 BF681641.1	0.0E+00 BE903372.1	0.0E+00 BF312552.1	1.				0.0E+00 BE698861.1	0.0E+00 BE698881.1		7669505 NT	7669505 NT	11024711 NT	
Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00 BE	0.0E+00 B	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+30	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 X51755.1	0.0E+00 X51755.1	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	
Expression Signal	2.57	5.52	2.04	2.04	63.21	4.08	3.25	37.53	2.93	6.35	1.83	1.93	1.93	2.32	2.32	5.68	5.68	3.22	6.15	6.15	3.02	3.02	20.38	1.98	1.98	31.56	38.47	36.47	34.29	
ORF SEQ ID NO:	28983			28390	26434	26006	29005	29006	29007				29017	29018	29019	28020	29021	29026	28034	29035	29036	29037		29062	29063	28066	29076	28077	29078	
Exan SEQ ID NO:		18694		18696	16272	15883	18710						18724	18725	18725	18727	18727		ı	18741	- 1		ł		18771	18775		ı	18787	
Probe SEQ ID NO:	8878	8883	8885	8882	8896	8900	8902	8903	8904	8915	8916	8916	8916	8917	8917	8919	8919	8924	8933	8933	8934	8934	8964	8962	8965	8969	8984	8981	8982	

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Table 4
Single Exon Probes Expressed in Heart

	_	_	_			_			_	_							В		4	41	, o a	4		4	•	Here	16.	B R.	d 25.
Top Hit Descriptor	HSB77E122 STRATAGENE Human skeletal muscle cDNA library, cat. #936215. Homo saplens cDNA clone 77E12	Human Chediak-Higashi syndrome protein short isoform (LYST) mRNA, complete cds	Homo sapiens of cardiac alpha-myosin heavy chain gene	601150023F1 NIH_MGC_19 Homo saplens cDNA clone IMAGE:3503020 5'	Homo saplens chromosome 21 segment HS21 C046	q917b12.x1 Soares_fetal_lung_NbHL19W Homo sapiens cDNA clone IMAGE:1739231 3'	Hamo sapiens gene for AF-6, complete cds	Hame sapiens chromosome 21 segment HS2/C046	Homo sapiens calcineurin binding protein 1 (KIAA0330), mRNA	Homo saplens anticodeant protein 1 (AOP1), nuclear gene encoding mitochondrial protein, mRNA	Homo sapiens glutathione S-transferase theta 2 (GSTT2) and glutathione S-transferase theta 1 (GSTT1)	genes, without ws DKFZb434K0819 r1 434 (synonym: htes3) Homo saniens cDNA cione DKFZn434K0819 5	Homo sapiens G-2 and S-phase expressed 1 (GTSE1), mRNA	DKFZp434G218_r1 434 (synonym: htes3) Horio sapiens cDNA clone DKFZp434G218 5	IL-BT030-271098-001 BT030 Homo sapiens cDNA	yv40e08.s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA done IMAGE:245222 3' similar to	SW POL_BAEVM P10272 POL POLYPROTEIN;	Homo sapiens adenylosuccinate lyase gene, complete cds	Homo sapiens T-cell lymphoma invasion and metastasis 1 (TIAM1) mRNA	Homo sepiens T-cell lymphoma invasion and metastasis 1 (TIAM1) mRNA	Homo sapiens nuclear factor of activated T-cells, cytopiacmic, calcineurin-dependent 2 (NFATC2), mRNA	Homo sapiens X-linked anhidrotic ectodermal dysplasia protein gene (EDA), exon 2 and flanking repeat rections	Homo sapiens low density tipoprotein-related protein 2 (LRP2), mRNA	hg31e06.X1 NCI_CGAP_GC6 Homo sapiens cDNA clone IMAGE:2847234 3' similar to contains Alu repetitive element contains element MER22 repetitive element.	RC8-BT0711-290300-011-D05 BT0711 Homo sapiens cDNA	Homo saplens somatostatin receptor subtype 3 (SSTR3) gene, 6' flanking region and partial cds	Human endogenous retrovirus, complete genome	an05h04.x1 Stratagene schizo brain S11 Homo sapiens cDNA clone IMAGE:1684759 3'	QV-BT065-020399-103 BT065 Homo capiens cDNA
Top Hit Database Source	EST HUMAN	N	NT	EST HUMAN	NT	EST_HUMAN	N	LN TN	١	NT	H	EST HUMAN	7	EST HUMAN	EST_HUMAN		EST HUMAN	LN.	LN	Į.	NT	IN	LN LN	EST HUMAN	EST HUMAN	NT	FN	EST_HUMAN	EST_HUMAN
Тф Hit Acession No.	F00884.1	U84744.1	220656.1	BE312542.1	AL163246.2	Al190993.1	AB011399.1	AL163246.2	11417862 NT	5802973 NT	A 5240788 4	AL041931.1	11418318 NT	AL046544.1	AI903497.1		N54484.1	AF106656.1	4507500 NT	4507500 NT	10092587 NT	AF003528.1	11430460 NT	AW590082.1	BE090210.1	AF068757.1	9635487 NT	AI204914.1	A1904646.1
Most Similar (Top) Hit BLAST E Value	0.0E+00	_	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	00730	_	-		0.0E+00		_		0.0E+00	0.0E+00	0.0E+00	0.0E+00	-	0.0E+00	_		0.0E+00	_	0.0E+00
Expression Signal	31.52	7.35	92.9	2.54	2.67	3.43	2.24	2.2	2.73	5.48	8	2.82	3.07	4.39	2.38	,	1.3	3.36	3.21	3.21	2.75	2.7	2.48	3.23	1.61	2.33	1.56	1.59	1.58
ORF SEQ ID NO:				24833							89036								20601				25183	25084					
Exan SEQ ID NO:	18792	18803	18805	19747					18849	18864	10563		ŀ	18910	19610		- [-1	- [10752	19612	10477	19412	19544	19595	19607	19092	i	19136
Probe SEQ ID NO:	8987	0008	8005	9017	9031	9033	9043	8062	9071	0606	67.23	9133	9158	9167	9180	2,000	8178	8233	9236	9236	9246	9276	9309	9370	9382	9426	9461	9498	8529

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WO 01/57274

Top Hit Descriptor	HTM1-654F HTM1 Homo sapiens cDNA	Homo sapiens calcineurin binding protein 1 (KIAA0330), mRNA	Homo sapiens calcineurin binding protein 1 (KIAA0330), mRNA	Homo sapiens caveolin-3 (CAV3) mRNA, complete cds	уобзеов.r1 Soares breast 3NbHBst Homo saplens cDNA clone IMAGE:182246 5' similar to gb:M64099 GAMMA-GLUTAMYLTRANSPEPTIDASE 5 PRECURSOR (HUMAN);	yoS9e08.r1 Soares breast 3NbHBst Homo sapiens cDNA clone IMAGE:182246 5' similar to gb:M64099 GAMMA-GLUTAMYLTRANSPEPTIDASE 5 PRECURSOR (HUMAN);	Human gamma-cytoplasmic actin (ACTGP9) pseudogene	Homo sapiens thyroid autoantigen 70kD (Ku antigen) (G22P1), mRNA	Homo sapiens thyroid autoantigen 70kD (Ku antigen) (G22P1), mRNA	TCBAP1E4468 Pediatric pre-B cell acute lymphoblastic leukemia Baylor-HGSC project=TCBA Homo sapiens	cDNA clone TCBAP4466	Hamo sapiens hypothetical protein FLJ10697 (FLJ10697), mRNA	Homo saplens hypothetical protein FLJ20454 (FLJ20454), mRNA	Homo sapiens G protein-coupled receptor 24 (GPR24), mRNA	Homo sapiens CST gene for cerebroside sulfcitransferase, exon 1, 2, 3, 4, 5	Homo saplens cleavage and polyadenylation specific factor 1, 160kD subunit (OPSF1), mRNA	Homo sapiens chromosome 21 segment HS21 C046	Homo saplens low density lipoprotein-related protein 2 (LRP2), mRNA	Homo sapiens DKFZp434P211 protein (DKFZP434P211), mRNA	Homo sapiens period (Drosophila) homolog 3 (PER3), mRNA	Human endogenous retrovirus pHE.1 (ERV9)	Homo saplens oxytocin receptor (OXTR), mRNA	RC1-HT0595-200400-012-f12 HT0595 Homo sepiens cDNA	DKFZp434J0618_r1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434J0618
Top Hit Database Source	EST_HUMAN			LN	EST_HUMAN	EST_HUMAN	TN				EST_HUMAN				TN		LN				IN		EST_HUMAN	EST_HUMAN
Top Hit Acession No.	BE439792.1	6912457 NT	6912457 NT	AF036365.1	H30132.1	H30132.1	D50659.1	11418189 NT	11418189 NT		BE246780.1	8922593 NT	11526291 NT	4885312 NT	AB029900.1	9558724 NT	AL163246.2	6806918 NT	7657020 NT	8567387 NT	X57147.1	11434874 NT	BE177449.1	AL048911.1
Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00.1	0.0E+00	0.05+00	0.0E+00	0.0E+00	_	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00
Expression Signal	1.68	1.98	1.98	2.33	2.87	2.87	32.21	3.99	3.99		5.21	1.64	2.39	3.19	2.21	1.5	2.79	1.41	2.13	2.42	1.51	1.29	1.56	1.28
ORF SEQ ID NO:		21724	21725	25267	21428	21427		25244	25245		25214	54829		24886		25233		20354		25177				
Exan SEQ ID NO:	19596	11841	11841	19161	11561	11561	19179	19181	19181	ľ	19239	15089	19249	16092	19269	19291	19757	10546	19364		19407	13621	19591	19431
Probe SEQ ID NO:	9540	9531	9551	9571	9584	8584	8597	8589	9599		3685	2696	8698	9721	9734	8773	9794	0086	9878	9913	8538	9946	9968	8971

CLAIMS

A spatially-addressable set of single exon nucleic-acid probes for measuring gene expression in a sample derived
 from human heart comprising a plurality single exon nucleic probes, said probes comprising any one of the nucleotide sequences set out in SEQ ID NOs: 1 - 9,980 or a complementary sequence, or a portion of such a sequence.

- 10 2. A spatially-addressable set of single exon nucleic acid probes as claimed in claim 1 wherein each of said plurality of probes is separately and addressably amplifiable.
- 3. A spatially-addressable set of single exon nucleic acid probes as claimed in claim 1 wherein each of said plurality of probes is separately and addressably isolatable from said plurality.
- 4. A spatially-addressable set of single exon nucleic acid 20 probes as claimed in any of claims 1 to 3 wherein said probes comprise any one of the nucleotide sequences set out in SEQ ID NOS.: 9,981 - 19,771.
- 5. A spatially-addressable set of single exon nucleic acid probes as claimed in any of claims 1 to 4, wherein each of said plurality of probes is amplifiable using at least one common primer.
- 6. A spatially-addressable set of single exon nucleic acid 30 probes as claimed in any of claims 1 to 5 wherein the set comprises between 50 - 20,000 single exon nucleic acid probes.
- 7. A spatially-addressable set of single exon nucleic acid probes as claimed in any of claims 1 to 6, wherein the

average length of the single exon nucleic acid probes is between 200 and 500 bp.

- 8. A spatially-addressable set of single exon nucleic acid probes as claimed in any of claims 1 to 7, wherein at least 50% of said single exon nucleic acid probes lack prokaryotic and bacteriophage vector sequence.
- 9. A spatially-addressable set of single exon nucleic acid probes as claimed in any of claims 1 to 8, wherein at least 50% of said single exon nucleic acid probes lack homopolymeric stretches of A or T.
- 10. A spatially-addressable set of single exon nucleic acid probes as claimed in any of claims 1 - 9 characterised in that said set of probes is addressably disposed upon a substrate.
- 11. A spatially-addressable set of single exon nucleic acid 20 probes as claimed in claim 10 wherein said substrate is selected from glass, amorphous silicon, crystalline silicon and plastic.
- 12. A microarray comprising a spatially addressable set of25 single exon nucleic acid probes as claimed in any of claims1 11.
- 13. A single exon nucleic acid probe for measuring human gene expression in a sample derived from human heart
 30 comprising a nucleotide sequence as set out in any of SEQ ID NOs.: 1 9,980 or a complementary sequence or a fragment thereof wherein said probe hybridizes at high stringency to a nucleic acid molecule expressed in the human heart.

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14. A single exon nucleic acid probe as claimed in claim 13 comprising a nucleotide sequence as set out in any of SEQ ID NOs.: 9,981 - 19,771 or a complementary sequence or a fragment thereof.

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- 15. A single exon nucleic acid probe for measuring human gene expression in a sample derived from human heart which is a nucleic acid molecule having a sequence encoding a peptide comprising a peptide sequence as set out in any of SEQ ID NOs.: 19,772 29,119, or a complementary sequence or a fragment thereof wherein said probe hybridizes at high stringency to a nucleic acid expressed in the human heart.
- 16. A single exon nucleic acid probe as claimed in any one of claims 13 to 15 wherein said single exon nucleic acid probe comprises between 15 and 25 contiguous nucleotides of said SEQ ID NO.
- 17. A single exon nucleic acid probe as claimed in any one of claims 13 to 15, wherein said probe is between 3 25 kb in length.
 - 18. A single exon nucleic acid probe as claimed in any one of claims 13 17, wherein said probe is DNA, RNA or PNA.

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- 19. A single exon nucleic acid probe as claimed in any one of claims 13 18, wherein said probe is detectably labeled.
- 30 20. A single exon nucleic acid probe as claimed in any one of claims 13 19, wherein said probe lacks prokaryotic and bacteriophage vector sequence.
- 21. A single exon nucleic acid probe as claimed in any one 35 of claims 13 - 20, wherein said probe lacks homopolymeric

stretches of A or T.

22. A method of measuring gene expression in a sample derived from human heart, comprising:

contacting the microarray of claim 12, with a first collection of detectably labeled nucleic acids, said first collection of nucleic acids derived from mRNA of human heart; and then

measuring the label detectably bound to each probe of said microarray.

23. A method of identifying exons in a eukaryotic genome, comprising:

algorithmically predicting at least one exon from genomic sequence of said eukaryote; and then detecting specific hybridization of detectably labeled nucleic acids to a single exon probe,

wherein said detectably labeled nucleic acids are derived from mRNA from the heart of said eukaryote, said probe is a single exon probe having a fragment identical in sequence to, or complementary in sequence to, said predicted exon, said probe is included within a microarray according to claim 12, and said fragment is selectively hybridizable at high stringency.

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24. A method of assigning exons to a single gene, comprising:

identifying a plurality of exons from genomic sequence according to the method of claim 23; and then

measuring the expression of each of said exons in a plurality of tissues and/or cell types using hybridization to single exon microarrays having a probe with said exon,

35 wherein a common pattern of expression of said exons in

said plurality of tissues and/or cell types indicates that the exons should be assigned to a single gene.

- 25. A nucleic acid sequence as set out in any of SEQ ID $5 ext{ NOs: } 1 19,771 ext{ which encodes a peptide.}$
 - 26. A peptide encoded by a sequence as set out in any of SEQ ID Nos: 1 19,771.
- 10 27. A peptide comprising a sequence as set out in any of SEQ ID Nos: 19,772 29,119.

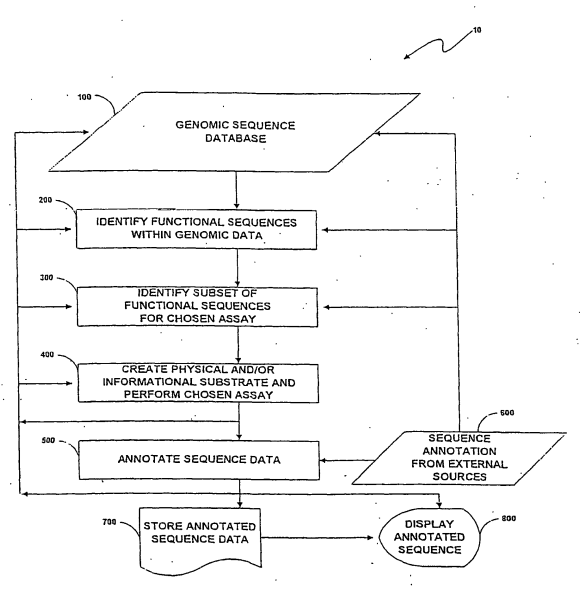


Fig. 1

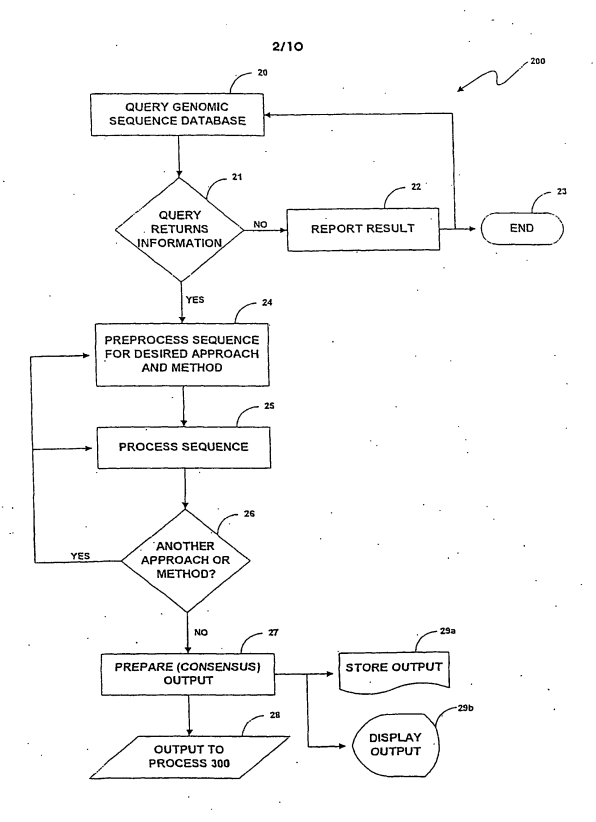


Fig. 2

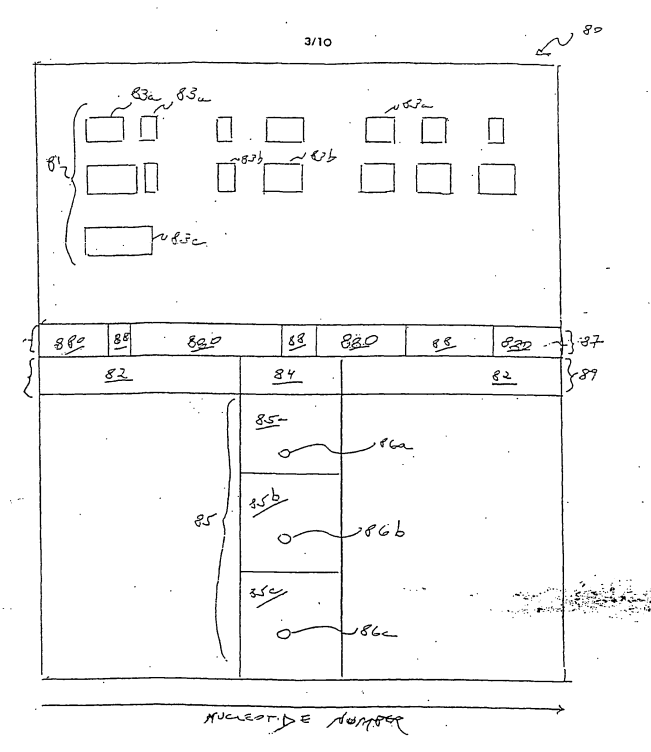


Fig. 3

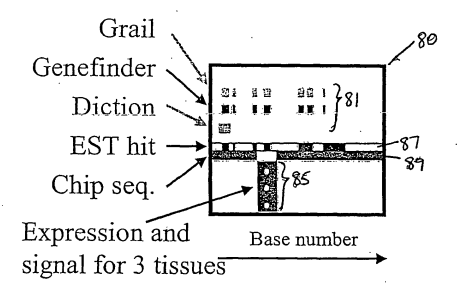


Fig. 4

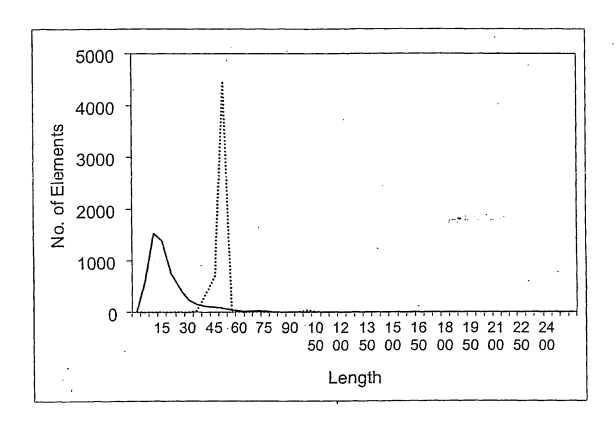


Fig. 5

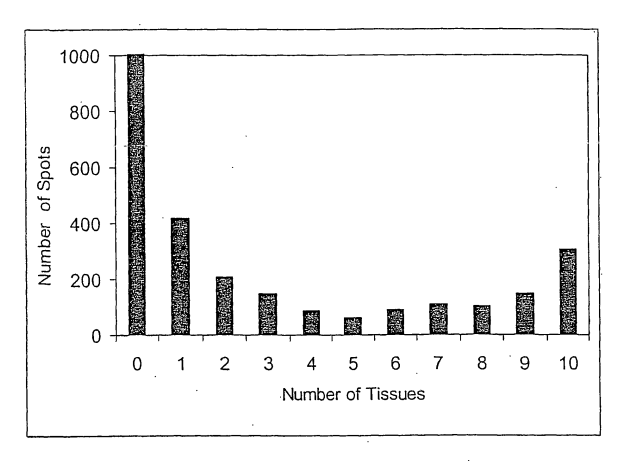
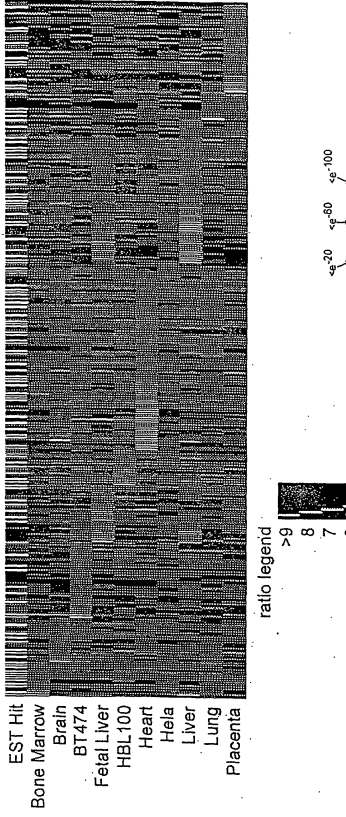


Fig. 6

7/10



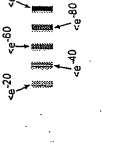


Fig. 7b

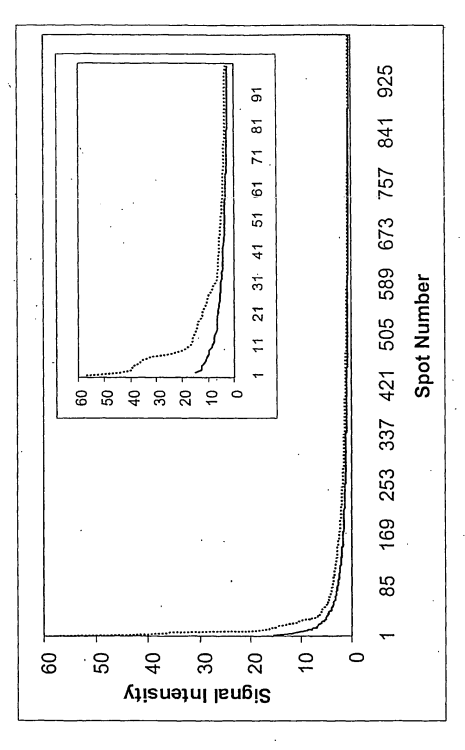
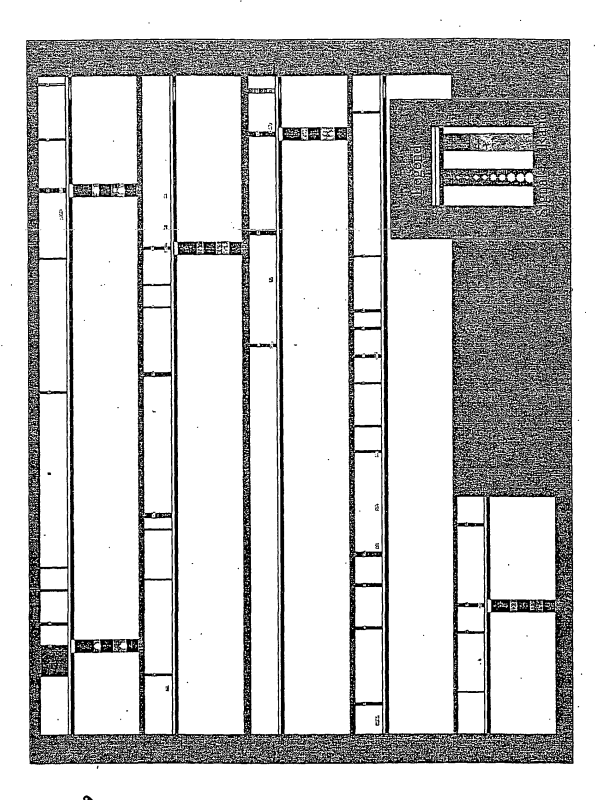
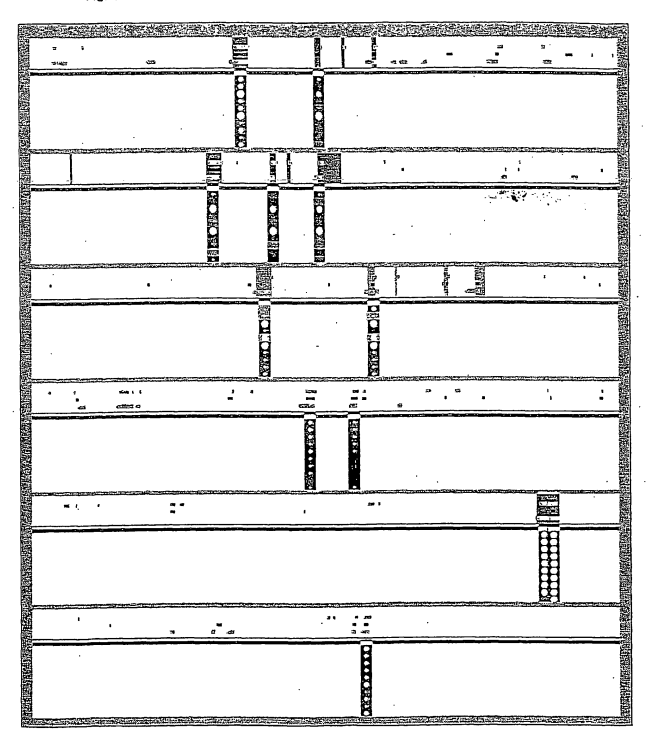


Fig. 8



<u>.</u>

Fig. 10



(19) World Intellectual Property Organization International Bureau





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-

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09/632,366	3 August 2000 (03.08.2000)	US
60/234,687	21 September 2000 (21.09.2000)	US
60/236,359	27 September 2000 (27.09.2000)	US
0024263.6	4 October 2000 (04.10.2000)	GB

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(72) Inventors; and

- (75) Inventors/Applicants (for US only): PENN, Sharron, G. [GB/US]; 617 South Delaware Street, San Mateo, CA 94402 (US). HANZEL, David, K. [US/US]; 968 Loma Verde Avenue, Palo Alto, CA 94303 (US). CHEN, Wensheng [CN/US]; 210 Easy Street #25, Mountain View, CA 94043 (US). RANK, David, R. [US/US]; 117 El Dorado Commons, Fremont, CA 94539 (US).
- (74) Agent: RONNING, Royal, N., Jr.; Amersham Pharmacia Biotech, Inc., 800 Centennial Avenue, P.O. Box 1327, Piscataway, NJ 08855 (US).

(81) Designated States (national): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW.

(84) Designated States (regional): ARIPO patent (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

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37274 A3

(54) Title: HUMAN GENOME-DERIVED SINGLE EXON NUCLEIC ACID PROBES USEFUL FOR ANALYSIS OF GENE EXPRESSION IN HUMAN HEART

(57) Abstract: A single exon nucleic acid microarray comprising a plurality of single exon nucleic acid probes for measuring gene expression in a sample derived from human heart is described. Also described are single exon nucleic acid probes expressed in the heart and their use in methods for detecting gene expression.

INTERNATIONAL SEARCH REPORT

ernational Application No PCT/US 01/00666

A. CLASSIFICATION OF SUBJECT MATTER IPC 7 C1201/68

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

SEQUENCE SEARCH, EPO-Internal, PAJ, WPI Data, BIOSIS

Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	DATABASE EBI 'Online! EMBL; Accession Number: AL049837, 11 May 1999 (1999-05-11) HEILIG ET AL.: "Human chromosome 14 DNA sequence BAC R-959A22 of library RPCI-11 from chromosome 14 of Homo sapiens (Human)" XP002182997	13-21, 25,27
,	the whole document/	1-12, 22-24,26

X Further documents are listed in the continuation of box C.	Patent family members are listed in annex.
Special categories of cited documents: A document defining the general state of the art which is not considered to be of particular relevance E earlier document but published on or after the international filing date L document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) O document referring to an oral disclosure, use, exhibition or other means P document published prior to the international filing date but later than the priority date claimed	"T" later document published after the international filling date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken atone "Y" document of particular relevance; the claimed invention cannot be considered to involve an invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art. "&" document member of the same patent family
Date of the actual completion of the international search	Date of mailing of the International search report
12 July 2002	0 7. 08. 2002
Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL – 2280 HV Rijswljk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Authorized officer Botz, J

INTERNATIONAL SEARCH REPORT

ternational Application No
PCT/US 01/00666

	Istion) DOCUMENTS CONSIDERED TO BE RELEVANT	Date and the state of the
Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	DATABASE EBI 'Online!	13-21,
^	EMBL;	25,27
	AA414703, 9 May 1997 (1997-05-09)],-
	MARRA ET AL.: "The WashU-HHMI Mouse EST	
	Project"	
	XP002205620	
Y	the whole document	1-12,
		22-24,26
X	DATABASE EBI 'Online!	15-21,27
	EMBL;	1
	Accession Number: 002711,	
	1 July 1997 (1997-07-01) BENIT ET AL.: "Cloning of a new murine	ļ
	endogenous retrovirus MuERV-L with strong	ļ
	similarity to the human HERV-L element and	
	with a gag coding sequence closely related	
	to the Fv1 restriction gene"	1
	XP002037954	
Υ	the whole document	1-12,
		22-24,26
		
Y	LIEW ET AL .: "A catalogue of genes in	1-27
	the cardiovascular system as identified by	
	expressed sequence tags"	Į
	PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES, USA,	}
	vol. 91, October 1994 (1994-10), pages	
	10645-10649, XP002185728	
	table 2	
Υ	LIPSHUTZ ET AL.: "High density synthetic	1–27
	oligonucleotide arrays"	
	NATURE GENETICS,	
	vol. 21, January 1999 (1999-01), pages	
	20-24, XP002115232	
	NEW YÓRK, NY, US the whole document	į
		j
A	DATSON ET AL.: "Scanning for genes in	Į
•	large genomic regions: cosmid based exon	į.
	trapping of multipleexpons in a single	
	product"	
	NUCLEIC ACIDS RESEARCH, OXFORD UNIVERSITY	
	PRESS,	
	vol. 24, no. 6, 1996, pages 1105-1111,	
	XP002081565	İ
	Surrey, GB	!
	the whole document	
A	WO 98 30722 A (MACK DAVID H.)	
n	16 July 1998 (1998-07-16)	
	page 63, paragraph 2	
	hade eat hai all ali	

International application No. PCT/US 01/00666

INTERNATIONAL SEARCH REPORT

Box I Obs rvations wher certain claims were found unsearchable (Continuation of item 1 of first she t)
This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2. X Claims Nos.: 1-12, 15-21 (partially not searched) because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful international Search can be carried out, specifically: See FURTHER INFORMATION sheet PCT/ISA/210
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This International Searching Authority found multiple inventions in this international application, as follows:
see additional sheet
As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. X As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.: 1-27 (partially)
4. No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark on Protest The additional search fees were accompanied by the applicant's protest. X No protest accompanied the payment of additional search fees.

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

Continuation of Box I.2

Claims Nos.: 1-12, 15-21 (partially not searched)

The following statements about the impossibility of performing a meaningful search according to Art. 17(2) PCT are made for the subject matter for which a search has been performed and identified as the first and second inventions in form 206 PCT.

Present claims 1-12 and 22-24 relate to an extremely large number of possible sets of nucleic acid probes comprising Seq. Id. 1 or 2 as well as microarrays comprising said sets. In fact, the claims contain so many possible permutations that a lack of clarity and conciseness within the meaning of Article 6 PCT arises to such an extent as to render a meaningful search of the claims impossible. Consequently, the search for the sets of probes comprising Seq. Id. 1 or 2 has been limited to the Seq. Id. as such.

Claims 1~3, 5, 6, 8-15 and 18-24 relate to portions or fragments of nucleic acids defined by Seq. Id. 1 or 2. The length or other similar characterizing features of the portions or fragments is not disclosed, bringing the total number of possible prior art sequences to exceptionally high numbers. The shorter the length, the higher the possibility that an overflow of, in principle unrelated, sequences are retrieved, making the establishment of a meaningful International Search Report impossible. For this reason the search has been limited to portions or fragments of Seq. Id. 1 or 2 having a significant minimum length and being supported by the description, namely at least 15 contiguous nucleotides (se claim 16).

Claims 15-21 relate to an extremely large number of nucleic acid probes. The probes are defined solely by their potential to code for peptide Seq. Id. 19780. However, due to the degeneracy of the genetic code, every peptide is potentially coded by an extremely high number of nucleic acid sequences. In fact, the claims contain so many potential nucleic acid sequences that a lack of clarity and conciseness within the meaning of Article 6 PCT arises to such an extent as to render a meaningful search over the whole scope of the claims impossible. The search has therefore been carried out for those parts of the claims which do appear to be clear and concise, namely the nucleic acid sequences disclosed in the application and identified as encoding the referred peptide in table 4 (Seq. Ids. 1 or 2 and 9989).

Likewise, claim 26, which refers to peptides encoded by Seq. Id. 1 or 2 and 9989, encompasses a high and undefined number of possible peptides. Besides three possible reading frames deriving from the encoding nucleic acid strand, as well as three additional reading frames deriving from the complementary nucleic acid strand, every possible fragment of these is being covered by the claim. This is due to the potential presence of stop codons within any of the six possible reading frames which can not be established a priori. Thus, claim 26 contains so many potential peptide sequences that a lack of clarity and conciseness within the meaning of Article 6 PCT arises to such an extent as to render a meaningful search

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

over the whole scope of the claim impossible. Consequently, the search has been carried out for those parts of the claim which do appear to be clear and concise, namely the peptide disclosed, identified by Seq. Id. 19780.

The applicant's attention is drawn to the fact that claims, or parts of claims, relating to inventions in respect of which no international search report has been established need not be the subject of an international preliminary examination (Rule 66.1(e) PCT). The applicant is advised that the EPO policy when acting as an International Preliminary Examining Authority is normally not to carry out a preliminary examination on matter which has not been searched. This is the case irrespective of whether or not the claims are amended following receipt of the search report or during any Chapter II procedure.

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

This International Searching Authority found multiple (groups of) inventions in this international application, as follows:

1. Claims: 1 - 27 (partially)

Invention number 1:

A nucleic acid probe comprising SEQ ID 1, complementary sequences or fragments thereof (in particular comprising Seq. Id. 9989). Spatially addressable sets of probes comprising said sequence, microarrays comprising said sets, a method for measuring gene expression, a method for identifying exons, a method for assigning exons to a single gene comprising the use of said arrays and peptide encoded by Seq. Id. 1 (in particular the one defined by Seq. Id. 19780).

2. Claims: 1 - 27 (partially)

Invention 2

A nucleic acid probe comprising SEQ ID 2, complementary sequences or fragments thereof (in particular comprising Seq. Id. 9989). Spatially addressable sets of probes comprising said sequence, microarrays comprising said sets, a method for measuring gene expression, a method for identifying exons, a method for assigning exons to a single gene comprising the use of said arrays and peptide encoded by Seq. Id. 2 (in particular the one defined by Seq. Id. 19780).

3. Claims: 1 - 27 (partially)

Inventions 3 - 9980

A nucleic acid probe comprising SEQ ID n (where n ranges from 2 - 9980 according to the invention number above), complementary sequences or fragments thereof, in particular comprising the SEQ ID no. which is listed in the column "Exon Seq. Id. no." in the same row that contains Seq. Id. n in table 4. Spatially addressable sets of probes comprising said sequence, microarrays comprising said sets, a method for measuring gene expression, a method for identifying exons, a method for assigning exons to a single gene comprising the use of said arrays and peptide encoded by Seq. Id. n, in particular the one defined by the Seq. Id. no. in the column "ORF Seq. Id. no." of the same row where Seq. Id. n is listed.

INTERNATIONAL SEARCH REPORT

Information on patent family members

rternational Application No

Patent document cited in search report	Publication date	Patent family member(s)	Publication date
WO 9830722 A	16-07-1998	AU 6035698 A EP 0973939 A1 JP 2001508303 T US 6303301 B1 WO 9830722 A1 US 2002028454 A1 US 2002039739 A1	03-08-1998 26-01-2000 26-06-2001 16-10-2001 16-07-1998 07-03-2002 04-04-2002

CORRECTED VERSION

(19) World Intellectual Property Organization International Bureau



(43) International Publication Date 9 August 2001 (09.08.2001)

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0024263.6	4 October 2000 (04.10.2000)	GB

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(72) Inventors; and

- (75) Inventors/Applicants (for US only): PENN, Sharron, G. [GB/US]; 617 South Delaware Street, San Mateo, CA 94402 (US). HANZEL, David, K. [US/US]; 968 Loma Verde Avenue, Palo Alto, CA 94303 (US). CHEN, Wensheng [CN/US]; 210 Easy Street #25, Mountain View, CA 94043 (US). RANK, David, R. [US/US]; 117 El Dorado Commons, Fremont, CA 94539 (US).
- (74) Agent: RONNING, Royal, N., Jr.; Amersham Pharmacia Biotech, Inc., 800 Centennial Avenue, P.O. Box 1327, Piscataway, NJ 08855 (US).

- (81) Designated States (national): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW.
- (84) Designated States (regional): ARIPO patent (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

Published:

- without international search report and to be republished upon receipt of that report
- with sequence listing part of description published separately in electronic form and available upon request from the International Bureau
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20 December 2001

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see PCT Gazette No. 51/2001 of 20 December 2001, Section II

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

7274 A2

(54) Title: HUMAN GENOME-DERIVED SINGLE EXON NUCLEIC ACID PROBES USEFUL FOR ANALYSIS OF GENE EXPRESSION IN HUMAN HEART

(57) Abstract: A single exon nucleic acid microarray comprising a plurality of single exon nucleic acid probes for measuring gene expression in a sample derived from human heart is described. Also described are single exon nucleic acid probes expressed in the heart and their use in methods for detecting gene expression.